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(54) POLYACETAL RESIN COMPOSITION

(57) A polyacetal resin composition comprises a opploacetal resin, and at least one carboxylic acid hydrazide selected from a seturated or unesturated longchain alliphatic carboxylic acid hydrazide, a situated or unesturated alloydic carboxylic acid hydrazide, a dimer acid or trimer acid hydrazide, and an oxycarboxylic acid hydrazide corresponding to each of these hydrazides. The proportion of the carboxylic acid hydrazide may be about 0.001 to 20 carts by welch fresilive to 100 parts by weight of the polyacetal resin. The polyacetal resin composition may further comprise at least one member selected from an antioxidant, a heat stabilizer, a processing stabilizer, a weather (light)-resistant stabilizer, a impact resistance improver, a slip-improving agent, a coloring agent, and a filler. With the use of such a resin composition, stability of a polyacetal resin is improved, and formatlehyde emission is inhibited.

Description

TECHNICAL FIELD

(0001) The present invertion relates to a polyacetal-series resin composition in which formaticetryde emission (or generation) is remarkably inhibited, extrusion properly and moldability are excellent, and blooming is suppressible; to a process of producing the same; and to a shaped (or molded) united formed from the resin composition.

BACKGROUND ART

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[0002] A polyacetal resin is excellent in mechanical proporty, fatigue resistance, friction or atrassion resistance, chemical realistance, and moltability. Therefore, the polyacetal resin has been widely utilized in various fields such as an automotive part, an electric or electronic device part, other precision machinery part, an erichtectural or pipeline part, an household utensil or cosmetic article part, or a medical device part. However, along with expansion or diversification in application, a polyacetal resin having higher qualifity has been demanded.

19003 Characteristics (or properties) required for the polyacetal resin include characteristics that mechanical strength in a process step such as an extunding step or a molding step is not deteriorated, that deposit to a metal mold (or mold deposit)) is not generated, that mechanical property under a long-term heating condition (heat aging) is adversely affected, and that incomplete (or defective) molding such as stifeva retract or void is not found in a shaped article. As one of the important factors responsible for such deterioration of strength or physical properties, and incomplete molding, the degradation of the oblymer upon heating is exemplified. In particular, the polyacetal resin is inherently unsable in an oxidative atmosphere sit are levelated temperature or in an action or sikaline environment because of its chemical structure. Therefore, the essential need that must be fulfilled for a polyacetal resin is that of insuring high thermal stability and minimal environment of cymarks of the control of the object of the control of processing and from shaped articles. Formsdelytic is chemically active and resety to be oxidized to formic acid to adversely affect the heat resistance of resin. In addition, when the resin is used as electric or electronic parts, formsdelptyde causes correction in metallic contacts or discolorion of the parts due to organic deposits, and contact errors occur. Furthermore, formsdelptyde itself pollutes the working environment in parts assembling as well as the living environment are around use of end products.

[004] In order to stabilize chamically active terminals, the following methods are known: for a homopolymer, a method of estatifying the terminal of the polymer by acetylation or other means; and for a copolymer, a method of copolymerizing troxane and a monomer having an adjacent cation bond (e.g., a cyclic ether or a cyclic formal), and then decomposing and removing unstable terminal sites to make the unstable terminal sites stable (or inactive) terminal sites. However, in a heading process, disavage (or fisson) decomposition also occurs in the main fainlain part of the polymer. Only the abovernentioned treatment is insufficient to prevent the polymer from such a decomposition, and practically, it is considered that addition of a stabilizer (e.g., an articolytaint, and other stabilizers) is essential for such inhibition.

[005] However, even in the case blending these stabilizers, it is difficult to completely inhibit decomposition (or degradation) of the polyacetal resin in practice, upon melt processing in an extruding step or a molding step for preparing a composition, the polyacetal resin undergoes an action of heat or exygen inside of explider of an extruder or a molding matchine, thereby generating formaldely/de from a decomposed main chain thereof or an insufficiently stabilized terminal hereof, as a result, working entrivonment is worsen in a extruding and molding process. Moreover, in the case carriering out molding for a long period, a finely powdered substance or a tar-like substance is deposited on a metal mold (mold deposit), thereby decreasing working efficiency. In addition, the mold deposit is one of the ultimate factors for deteriorating the surface condition of the shaped article. Further, the polymer decomposition causes deterioration in mechanical strength of the resin, and discoloration thereof. From such a viewpoint, a good deal of effort is continued for establishing more effective stabilizing formulation (or receipt about the polypocal resin.

[0006] As the antioxidant added to the polyacetal resin, a phenol-series (phenolic) compound having sterio hindrence (hindered phenol), and an amine compound having sterio hindrence (hindered amine) have been known. As other stabilizers, neelamine, an alkial metal hydroxid, an alkaline eath metal hydroxide, and an organic or inorganic acid sat have been known. Moreover, antioxidants are generally used in combination with other stabilizers. However, even when such an additive(s) is/are used, it is difficult to Inhibit formaldehyde emission (or generation) from a shaped encle of the polyacetal resin.

[0007] US Patent No. 3152/01 (Patent Document 1) discloses a composition comprising a polyacetal core in the rand a discriboxylic acid dihydrazide (e.g., an aliphatic discriboxylic acid dihydrazide), attouch use of such a short-chain aliphatic carboxylic acid dihydrazide, attouch use of such a short-chain aliphatic carboxylic acid dihydrazide improves heat stability at some level thereby inhibiting emission of formatideryde, such a composition is low in formability (or moldability). Therefore, mold deposit occurs, or the carboxylic acid hydrazide biedes out of a shaped article formad with the composition. Moreover, this document concretely fails to disclose what kind of alleyoic olicatroxylic acid dhydrazide improves heat stability of the resh composition.

[Patent Document 1] US Patent No. 3152101 (the first and the third columns)

DISCLOSURE OF THE INVENTION

PROBLEMS TO BE SOLVED BY THE INVENTION

[0008] It is therefore an object of the present invention to provide a polyacetal resin composition capable of improving heat (or thermal) stability of a polyacetal resin, and melt stability of a polyacetal resin in an extruding step or a molding step; and a process of producing the same; as well as an article as molded (or shaped) therefore.

[0003] It is another object of the present invention to provide a polyacetal resin composition conducive to a marked inhibition of formal dehyde emission with a small amount of an additive added, and improvement in working environment; and a process of producing the same; as well as an article as molded (or shaped) thorefrom.

[0010] It is still another object of the present invention to provide a polyacetal reein composition which is adapted for inhibiting emission of formaldehyde even under severe conditions to suppress deposition of decomposition products on the mod. blooming or bleeding of the decomposition products from a shaped article and themal deterioration of the stride, and which contributes to upgrading quality of the shaped article and improves the moldability; and a process of producing the same; as well as an article as molided (or shaped) therefrom.

[0011] It is further object of the present invention to provide a polyacetal resin composition in which an amount of formatlethyde emission from a polyacetal resin and a shaped activel teneform is inhibited to a significantly low fuel, and physical property such as weather (light)-resistant stability, impact resistance, or sliding property is improved; and anotice as motived (or shaped) therefrom.

MEANS TO SOLVE THE PROBLEMS

5 [0012] The inventor of the present invention made intensive studies or searches on a series of carboxylio acid hydrazide compounds regarding a stabilizer for a polyacetal resin to achieve the above objects and finally found that a specific acroscylic acid hydrazide compound significently inhibits formaldehyde emission from a shaped article of a polyacetal resin, and brings about an excellent modability and lowing of bleeding out of such a shaped article. The present invention was accomplished based on the above findings.

(0013) That is, the polyacetal resin composition of the present invention comprises a polyacetal resin and a carboxylic acid inydrazide. The carboxylic acid inydrazide comprises at least one member selected from the group consisting of a saturated or unsaturated or insert present production and produced and produced

6 be about 0.001 to 20 parts by weight relative to 100 parts by weight of the polyacetal resin. [0014] The polyacetal resin composition may further comprise at least one member selected from the group consisting of an antioxidant, a heat stabilizer, a processing stabilizer, a weather ([ght])-resistant stabilizer, an impact resistance improver, a stip-improving agent, a coloring agent, and a filler. The antioxidant, the processing stabilizer, the heat stabilizer, and the weather ([inith-resistant stabilizer may be substantially free from an intramolocular ester bond, incidentally, the resin composition of the present invention may improve in heat stability without substantially containing a phosphorus-containing fame retardant.

and an aliphatic oxyC₁₆₋₄₆dicarboxylic acid mono- or dihydrazide). The proportion of the carboxylic acid hydrazide may

[015] The present invention also includes a process for producing a polyacetal resin composition, which comprises mixing a polyacetal resin with the carboxytic acid hydrazide, wherein the resin composition is prepared by using an extruder, and feeding at least the carboxytic acid hydrazide through a side feed port of the extruder. In addition, the present invention further includes a shaped article formed from the polyacetal resin composition. The shaped article may be an automotive part, an electric orelectronic device part (an electric and/or-electronic device part, an architectual or pipeline part (an architectural and/or pipeline part (an architectural and/or pipeline part), and process the process of the proc

EFFECTS OF THE INVENTION

[0016] According to the present invention, addition of a specific carboxylic acid hydrazide, (a long-chain aliphatic carboxylic acid hydrazide, an alicyclic carboxylic acid hydrazide, an alicyclic carboxylic acid hydrazide, and environ or triver acid hydrazide, an oxycarboxylic acid hydrazide, by the polyacetal resin, and melt stability of the polyacetal resin in an extruding process or a molding process. Moreover, acidion of only a small amount of the carboxylic acid hydrazide significantly ensure to inhibit formakelynde general carboxylic acid hydrazide significantly ensure to inhibit formakelynde process. Moreover, acidion of other services of the process of the process

DETAILED DESCRIPTION OF THE INVENTION

20 [0017] The resin composition of the present invention comprises a polyacetal resin, and a specific carboxylic acid hydrazide.

[0018] (Polyacetal resin)

The polyacetal resin is a macromiscular compound containing oxymethylene group (-OCH₂-) as a predominant constituent unit and includes polyacetal homopolymens or polyoxymethylenes (e.g., trade name "Derlin", manufactured by DuPont, U.S.A.; trade name "Tearea 4010", manufactured by Asani Kasei Corp., etc.) and polyacetal copolymers comprising an oxymetrylene unit and a componer unit (e.g., trade name "Duracon", me-ifactured by Polyleistics Co., Ltd.). Referring to such coopylines, the componer unit includes oxysaltypine unit: af about 2 to 6 carbon atoms (preferably about 2 to 4 carbon atoms), for example, oxyethylene (-OCH₂CH₂-), oxyprocylene, and oxysteramethylene units. The proportion of such componer unit may be small and, for example, can be selected from the range of about 0.01 to 20 mole %, preferably about 0.03 to 15 mole % (e.g., 0.05 to 10 mole %), and more preferably about 0.1 to 10 mole %, relative to the whole polyacetal resin, the whole monomer units constituting the polyacetal resin.

[0019] The polyscetal copolymer may be, for example, a copolymer containing two components, a terpolymer containing three components and so on. The polyacetal copolymer may be also a random copolymer, a block copolymer (e.g., copolymers described in Japanese Petert Publication No. 2430/71990 (P.2-24307B), manufactured by Asshi Kasel Corp., trade name Tenac LA*, "Tenac LM*), or a graft copolymer. Moreover, the polyacetal resin may be linear or branched, and may have a crosalinked structure. In addition, the end (or terminal) groups of the polyacetal resin may nave been stabilized by esterification with a carboxylic acid such as acetic acid or propionic acid, or an anhydride thereof, urethanation with an isocyanate compound, or etherfication. There is no particular restriction as to the molecular weight of the polyacetal, provided it can be only melt-imidied. There is no particular restriction as to the molecular weight of the polyacetal resin, and, for example, the weight average molecular weight also are to grant or example, the weight average molecular weight also are to grant or example, the weight average molecular weight also are to grant or example, the weight average

[0020] The polyacetal resin can be, for example, produced by polymerizing an aidehyde such as formaldehyde, paraformaldehyde, or acetaldehyde, or acyclic ether or cyclic formal such as trioxane, ethylene oxide, propylene oxide,
butlylene oxide, syrene oxide, cyclohexane oxide, gyclohexane oxide, 13-dioxane, 3-diehydene glycol
formal, or 1,4-butanediol formal. Further, as a copolymerizable component, an alkyl or anyiglycidyl ether (e.g., methylglycidyl ether, ethylgylcidyl ether, phenylgylcidyl ether, and naphthylgycidyl ether, an alkyl or anyiglycidyl ether, and be employed.
As a likyl or anyi glycidyl alcohol, a cyclic aster (e.g., β-propiolactone), or a vinyl compound (e.g., styrene, and vinyl ether)
can be employed.

50 [0021] (Carboxylic acid hydrazide)

The characteristic of the present invention resides in addition of a specific carboxylic acid hydrazide and thereby remarkably improving processing stability of a polyacetal resin and remarkably inhibiting generation of formaldehyde. [0022] The carboxylic acid hydrazide may include a saturated or unsaturated long-tehia leibhetic carboxylic acid hydrazide, a saturated or unsaturated alcyclic carboxylic acid hydrazide, a dimer acid or trimer acid hydrazide, and an oxycarboxylic acid hydrazide corresponding to each of these carboxylic acid hydrazides. The carboxylic acid constituting the carboxylic acid hydrazide may be a monocarboxylic acid, or may be a polycarboxylic acid (e.g., a discriboxylic acid and a ticarboxylic acid). The carboxylic acid hydrazide may include a monocarboxylic acid monohydrazide, a polycarboxylic acid mono- or polyhydrazide (e.g., a discriboxylic acid monohydrazide, and a discriboxylic acid displazation.

[9023] Examples of the long-chain aliphatic carboxylic acid hydrazide may include a saturated for unsaturated Co₁₄₋₀₄ aliphatic carboxylic acid hydrazide (for example, a monocarboxylic acid monocarboxylic carboxylic acid monocarboxylic acid monocarboxylic acid monocarboxylic acid monocarboxylic acid monocarboxylic acid monocarboxylic acid hydrazide, incentical hydrazide, acid hydrazide, monocarboxylic acid hydrazide, accerd acid hydrazide, electrostated acid hydrazide, ac

[2024] Among them, the preferred one includes a saturated or unsaturated alliphatic C₁₆₋₄₀carboxylic acid hydrazide (e.g., a monocarboxylic acid monohydrazide, and a dicarboxylic acid mono- or dhightazide), pertibulanty, emonohydrazide of a C₁₆₋₄₀casturated aliphatic carboxylic acid, such as behenic acid hydrazide or montains acid hydrazide; a dhightazide of a C₁₆₋₄₀casturated aliphatic carboxylic acid, such as 12-eioosanedioicacid dihydrazide; and a C₁₆₋₄₀casturated aliphatic carboxylic acid, such as 12-eioosanedioicacid dihydrazide; and a C₁₆₋₄₀casturated aliphatic carboxylic acid on ydrazide such as 8,12-eioosaldenedioicacid dihydrazide. Includentally, the details of processes for producing elocsanedioic acid dihydrazide. Includentally, the details of processes for producing elocsanedioic acid dihydrazide. Includentally, the details of processes for producing elocsanedioic acid dihydrazide. Includentally, the details of processes for producing elocsanedioic acid dihydrazide.

1025] The overactiony's each phystratic corresponding to the long-chain alighatic carboxylic acid hydrazide may have at least one hydroxyl group per one molecule. The number of hydroxyl groups is not particularly limited to a specific one, and may be, for example, a bout 1 to 6, preferably about 1 to 4, and more preferably about 1 to 3. Such an oxycarboxylic acid hydrazide and may help of the may include, for example, a seturated or unsaturated alighatic oxyC₁₆₋₄₀catboxylic acid hydrazide such as a staturated affighatic oxycarboxylic acid hydrazide such as a staturated affighatic oxycarboxylic acid hydrazide such as a staturated affighatic oxycarboxylic acid mono-hydrazide such as a staturated affighatic oxycarboxylic acid, such as a hydroxystearic acid hydrazide, and e. 10-dihydroxycarboxylic acid mono- or polyhydrazide, and hydroxydexadecan-dolo acid mono- or dihydrazide, and hydroxydexadecan-dolo acid mono- or dihydrazide, and hydroxydexadecan-dolo acid hydrazide, and hydroxydexadecan-dolo acid hydrazide, alientific acid hydrazide, alientific acid hydrazide, and hydroxydexadecan-dolo acid hydrazide, alientific acid hydrazide, and hydroxydexadecan-dolo acid hydrazide, alientific acid hydrazide, and hydroxydexadecan-dolo acid hydrazide, alientific acid hydrazide, and alientific acid hydrazide, and alientific acid hydrazide, or phelionic acid hydrazide, and an alientific acid hydrazide, and a sydroxydicarboxylic acid mono- or dihydrazide, and an alientific acydrazide, and an alientific acydrazide acid hydrazide, and an alientific acid mono- or dihydrazide, and an alientific acid hydrazide, and an alientific acid hydrazide.

[0026] Among them, a saturated or unsaturated allphatic $cxyC_{16,40}$ carboxylic acid hydrazide [in particular, a saturated or unsaturated allphatic $cxyC_{16,40}$ carboxylic acid hydrazide [in particular, a saturated or unsaturated allphatic $cxyC_{16,40}$ carboxylic acid hydrazide) (particularly 12-hydroxystearic acid hydrazide), and the control of the control

[0027] Among the alicyclic carboxylic acid hydrazides, as the saturated alicyclic carboxylic acid hydrazide, there may be mentioned a saturated C_{6.40}alicyclic carboxylic acid hydrazide such as a monocarboxylic acid monohydrazide (for example, a cycloalkanecarboxylic acid hydrazide such as a cyclopentanecarboxylic acid hydrazide, a cyclohexanecarboxylic acid hydrazide, a cycloheptanecarboxylic acid hydrazide, a cyclooctanecarboxylic acid hydrazide, a cyclononanecarboxylic acid hydrazide, a cyclodecanecarboxylic acid hydrazide, a cycloundecanecarboxylic acid hydrazide, or a cyclododecanecarboxylic acid hydrazide; a polycycloalkanecarboxylic acid hydrazide such as a norbornanecarboxylic acid hydrazide, or an adamantanecarboxylic acid hydrazide; and a cycloalkyl-alkanecarboxylic acid hydrazide such as cyclohexylecetic acid hydrazide, or cyclohexylbutyric acid hydrazide), or a polycarboxylic acid mono- or polyhydrazide for example, a di- to tetracarboxylic acid mono- to tetrahydrazide corresponding to each of the monocarboxylic acid monohydrazides (e.g., a dicarboxylic acid mono- or dihydrazide, a tricarboxylic acid mono- to trihydrazide, and a tetracarboxylic acid mono- to tetrahydrazide), for example, a cycloalkane di- or tricarboxylic acid hydrazide such as a 1,4cyclohexanedicarboxylic acid mono- or dihydrazide, or 1,3,5-cyclohexanetricarboxylic acid mono-to trihydrazide (e.g., a cycloalkanedicarboxylic acid mono-or dihydrazide, and a cycloalkanetricarboxylic acid mono-to trihydrazide); a polycycloalkanedicarboxylic acid mono-or dihydrazide; a carboxyalkylcycloalkanecarboxylic acid monohydrazide, a hydrazinocarbonylalkylcycloaikanemonocarboxylic acid monohydrazide, a hydrazinocarbonylalkylcycloaikanecarboxylic acid, and a di/hydrazinocarbonylalky/)cycloalkane).

[0028] Among the alloydic carboxylic acid hydrazides, the unsaturated alloydic carboxylic acid hydrazide may include an unsaturated C₆₋₄₀elicyclic carboxylic acid hydrazide such as amonocarboxylic acid monohydrazide(e.g., e cyclakenecarboxylic acid hydrazide, e cyclohexenecarboxylic acid hydrazide, e cyclohexenecarboxylic acid hydrazide, except such as a cyclopentenecarboxylic acid hydrazide such as a cyclopentenecarboxylic acid hydrazide, a cyclopentenecarboxylic acid hydrazide such as a cyclopentenecarboxylic acid hydrazide, a cyclopentenecarboxylic acid hydrazide such as a cyclopentenecarboxylic acid hydrazide acid hydraz

hydrazide, a cycloheptenecarboxylic acid hydrazide, or a cyclooctenecarboxylic acid hydrazide; a polycycloalkenecarboxviic acid hydrazide such as norbornenecarboxylic acid hydrazide; a cycloalkenylakanecarboxylic acid hydrazide such as hydrocarpic acid hydrazide, chaulmoogric acid hydrazide, or gorlic acid hydrazide), a polycarboxylic acid monoor polyhydrazide [for example, a di- to tetracerboxylic acid mono- to tetra hydrazide corresponding to each of the abovementioned monocarboxylic acid monohydrazides (e.g., a dicarboxylic acid mono- or dihydrazide (e.g., a cycloalkenedicarboxylic acid mono- or dihydrazide), a tricarboxylic acid mono- to trihydrazide (e.g., a cycloalkenetricarboxylic acid mono- to trihydrazide), and a tetracarboxylic acid mono- to tetrahydrazide (e.g., a cycloalkenetetracarboxylic acid monoto tetrahydrazide))]; or a trimer acid (e.g., a trimer acid of an unsaturated aliphatic carboxylic acid, such as a cyclic unsaturated linoleic trimer acid, or a cyclic unsaturated linolenic trimer acid) mono- to trihydrazide). Incidentally, the number of carbon-carbon unsaturated bonds in the unsaturated alicyclic carboxylic acid constituting the unsaturated alicyclic carboxylic acid hydrazide is not particularly limited to a specific one, and may be selected, depending on the member of an alloyolic ring thereof, from the range that the compound can maintain non-aromatic property. For example, the number of carbon-carbon unsaturated bonds may be about 1 to 6, preferably about 1 to 3, and more preferably 1 or 2. [0029] Incidentally, a saturated or unsaturated alloyolic carboxylic acid hydrazide having a polycyclic structure (e.g., a polycycloalkanecarboxylic acid hydrazide, and a polycycloalkenecarboxylic acid hydrazide) may be obtained by hydrazidation of a Diels-Alder reaction product between an unsaturated alicyclic compound having 1,3-diene structure (e.g., a cycloalkadi- to tetra-ene such as cyclopentadiene, dicyclopentadiene, cyclohexadiene, or cyclopetadi- to tetraene) and an olefin compound [e.g., an alkyl (meth)acrylate, and maleic anhydride].

[0030] Among these alicyclic carboxylic acids, a saturated or unsaturated C_{e ze}alicyclic carboxylic acid hydrazide such as a cyclohexanedicarboxylic acid mono- or hydrazide), for example, an alicyclic mono- or dicarboxylic acid mono- or differencylic a

[0031] The alloyelic oxycarboxylic acid hydrazide may include an oxycarboxylic acid hydrazide corresponding to each of the above-mentioned alloyelic carboxylic acid hydrazides (e.g., a saturated or unsaturated alloyelic Q_{adjO} oxycarboxylic acid hydrazides (e.g., a saturated or unsaturated alloyelic Q_{adjO} oxycarboxylic acid south a second production of the south as a second product of the second product of t

5 hydroxycyclohexanecarboxylic acid hydrazide, a hydroxycyclohexanedicarboxylic acid hydrazide, or a hydroxycyc enylcarboxylic acid hydrazide (e.g., a hydroxyalicyclic monocarboxylic acid monorhydrazide, and a hydroxyalicy silcarboxylic acid mono- or dihydrazide).

[0032] It is sufficient that the alloyofic oxycarboxylic acid hydrazide has at least one hydroxyl group per one molecule thereof. The number of hydroxyl groups is not particularly limited to a specific one, and may be, for example, about 1 to 6, preferably about 1 to 7, and more preferably about 1 to 7.

[0033] The dimer acid or trimer acid hydrazide may be any of linear and cyclic one, and may be either a saturated or an unsaturated one. The dimer acid hydrazide or trimer acid hydrazide may holude a dimer acid mono- or polyhydrazide (for example, a dimer acid mono- or third read of mono- or polyhydrazide (for example, a dimer acid mono- or polyhydrazide), and there acid mono- or polyhydrazide (for example, a trimer acid mono- to texathydrazide (particularly, a trimer acid mono- to trihydrazide), and othere.

[0034] As the dimer acid or trimer acid constituting the dimer acid hydrazide or trimer acid hydrazide, there may be mentioned a dimer or trimer of an unsaturated aliphatic carboxylic acid (for example, an unsaturated C₁₀₋₀₃aiphatic carboxylic acid such as an unsaturated monocarboxylic acid (such as oleic acid, linoletc acid, or linolenic acid), and in addition an unsaturated dicarboxylic acid.

20 [0035] Examples of the dimar acid hydrazide may include a dimer acid hydrazide (e.g., a. A Cookdimer acid hydrazide), for example, a linear saturated or unsaturated dimer acid hydrazide such as a linear saturated (or hydrogenated) dimer acid hydrazide (e.g., a. linear saturated dimer acid mono- or dihydrazide), or a linear unsaturated dimer acid hydrazide (e.g., a.g. linear linear linear linear acid mono- or dihydrazide), a, a cyclic saturated or unsaturated dimer acid hydrazide (e.g., a.g. a cyclic saturated dimer acid mono- or or dihydrazide), or a cyclic unsaturated dimer acid mono- or dihydrazide), a cyclic unsaturated lineale dimer acid mono- or dihydrazide). As the trimer acid hydrazide, there may be mentioned a linear or cyclic trimer acid hydrazide corresponding to each of the dimer acid hydrazide (e.g., a.g. ox)einer acid nono- to trihydrazide), for example, all corresponding to each of the dimer acid hydrazide (e.g., a.g. ox)einer acid nono- to trihydrazide), for example, all corresponding to each of the dimer acid hydrazide (e.g., a.g. ox)einer acid hydrazide dihydrazide, acid hydrazide (e.g., a.g. ox)einer acid hydrazide (hydrazide).

[0036] Moreover, the dimer acid or trimer acid hydrazide may be an oxydimer acid or oxytrimer acid hydrazide having a hydroxyl group. Incidentally, in the unsaturated dimer acid hydrazide or unsaturated trimer acid hydrazide, the number of carbon-carbon unsaturated bonds is not particularly limited to a specific one, and may be, for example, about 1 to 6, preferably about 1 to 4, and more preferably 1 or 2.

[0037] Among these dimer acid or trimer acid hydrazides (a saturated or unsaturated dimer acid hydrazide, and a saturated or unsaturated trimer acid hydrazide), particularly, a linear C_{20-60} dimer acid mono- or dihydrazide (e.g., a

linear inoleic dimer acid hydrazide, and a linear inolenic dimer acid hydrazide), a cyclic C_{20-49} dimer acid mono- or dihydrazide (e.g., a cyclic linoleic dimer acid hydrazide), a linear C_{29-69} firmer acid mono- to trihydrazide, a cyclic C_{20-69} firmer acid mono- to trihydrazide (e.g., a cyclic linoleic trimer acid hydrazide), and others are preterred. Amog them, at least one member selected from the group consisting of montanic acid hydrazide, acid calcid dihydrazide, 8,12-closeadenediolc acid dihydrazide, 12-hydroxystearic acid hydrazide, 1,4-cyclohexanedicarroxylic acid dihydrazide, acid calcid dihydrazide, acid calcid dihydrazide, acid notecid dimer acid calcid calcid dihydrazide, acid calcid calcid dihydrazide, acid calcid calcid dihydrazide, acid calcid calcid dihydrazide is preterred.

[0038] These carboxylic acid hydrazides may be used singly or in combination.

[0039] Addition of an only small amount of the carboxylic acid hydrazide to a polyacetal resin brings about stabilizing effects far superior to conventional stabilizers, and thus obtained polyacetal resin composition is excellent in mobilizing (flame retardancy, mold deposit), Moreover, bleeding out (or blooming property) of such an additive from a shaped article formed from the polyacetal resin composition can be significantly improved.

[0040] The proportion of the carboxylic acid hydrazide may be selected from, for example, about 0.001 to 20 parts by weight, preferably about 0.002 to 10 parts by weight (e.g., about 0.002 to 5 parts by weight), and more preferably 0.003 to 3 parts by weight, relative to 100 parts by weight of the polyacetal resin. In particular, even when the proportion is about 0.005 to 2 parts by weight, generation of formal dehyde can be remarkably inhibited. In the case where the proportion of the carboxylic acid hydrazide is too low, it is difficult to effectively reduce the amount of formaldehyde emission. In the case where the proportion is too high, there is a possibility that moldability or mechanical strength is deteriorated. [0041] The carboxylic acid hydrazide can impart significant stability and processing stability to the polyacetal resin even when the carboxyllc acid hydrazide is used alone. Besides, the carboxylic acid hydrazide may be used in combination with at least one member selected from the group consisting of an antioxidant, a processing stabilizer, a heat stabilizer. a weather (light)-resistant stabilizer, an impact resistance improver, a slip-improving agent, a coloring agent, and a filler. [0042] Incidentally, many of the stabilizers (the antioxidant, the processing stabilizer, the heat stabilizer, and the weather (light)-resistant stabilizer) have an ester bond [-C(=O)O-] as a structural unit in a molecule thereof. The carboxylic actic hydrazide can stabilize the polyacetal resin even in the case of being used in combination with the stabilizer having such an ester bond. However, since the carboxylic acid hydrazide potentially has reactivity to an ester bond, the preferred stabilizer is a compound free from an ester bond as a structural unit in a molecule thereof in order to have an effect for inhibiting formaidehyde emission in a smaller amount of the carboxylic acid hydrazide.

[0043] (Antioxidant)

The antioxident may include a hindered phenof-series compound, and a hindered amine-series compound, and others. [0044] The hindered phenol-series compound may include a conventional phenol-series antioxidant or stabilizer, for example, a monocyclic hindered phenolic compound (e.g., 2,6-di-t-butyl-p-cresol), a polycyclic hindered phenolic compound in which rings are connected or bonded to each other through a hydrocarbon group or a group containing a sulfur atom [e.g., a C1,10 alky lene-bis to tetrakis(t-butylphenol) such as 2,2'-methylenebis(4-methyl-6-t-butylphenol), 4,4'-methvienebis(2.6-di-t-buty/iphenol) or 1,1,3-tris(2-methyl-4-hydroxy-5-t-buty/iphenyl)butane; a C2,10alkenylene or dienylenebis to tetrakis(t-butylphenol) such as 4,4'-butylidenebis(3-methyl-6-t-butylphenol); a C₆₋₂₀arylene or aralkylene-bis to tetrakis(t-butylphenol) such as 1,3,5-trimethyl-2,4,6-tris(3,5-di-t-butyl-4-hydroxybenzyl)benzene; and a bis(t-butylphenot) in which t-butylphenol groups are connected or bonded to each other through a group having a sulfur atom, for example, 4,4'-thiobis(3-methyl-6-t-butylphenol)], a hindered phenolic compound having an ester group or an amide group [e.g., a t-butylphenol having a C2,10alkylenecarbonyloxy group, exemplified by n-octadecyl-3-(4'-hydroxy-3'.5'-dit-butylphenyl)propionate or n-octadecyl-2-(4'-hydroxy-3',5'-di-t-butylphenyl)propionate; a bis to tetrakis(t-butylphenyl) In which t-butylphenol groups are connected or bonded to each other through a polyol ester of a fatty acid, exemplified bv 1.6-hexanedioi-bis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionate], triethylene glycol-bis[3-(3-t-butyl-5-methyl-4-hydroxyphenyl)propionate], droxyphenyl)propionate] or pentaerythritol tetrakis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionate]; a bis to tetrakis(tbutylphenol) having a heterocyclic group and a C2-10 alkylenecarbonyloxy group, exemplified by 3,9-bis[2-(3-t-butyl-4-hydroxy-5-methylphenyl)propionyloxy]-1,1-dimethylethyl]-2,4,8,10-tetraoxaspiro[5.5]undecane; a t-alkylphenol (e.g., t-butylphenol, and t-pentylphenol) having a Co. malkenylcarbonyloxy group, exemplified by 2-t-butyl-6-(3'-t-butyl-5'-methyi-2'-hydroxybenzyi)-4-methylphenylacrylate or 2-f1-(2-hydroxy-3,5-di-t-pentylphenyl)ethyll-4,6-di-t-pentylphenylacrviate; a hindered phenolic compound having a phosphonic ester group, exemplified by di-n-octadecyl-3.5-di-t-butyl-4hydroxybenzylphosphona te; a hindered phenolic compound having an amide unit, exemplified by N,N'-hexamethylenebis(3,5-di-t-butyl-4-hydroxy-dihydrocinnamamide), N,N'-ethylenebis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionamide], N.N'-tetramethylenebis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionamide], N,N'-hexamethylenebis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionamide], N,N'-hexamethylenebis[3-(3,5-di-t-butyl-4-hydroxyphenylbutyl-4-hydroxyphenyllpropionamidel. N.N'-ethylenebis[3-(3-t-butyl-5-methyl-4-hydroxyphenyllpropionamidel. N.N'hexamethylenebis[3-(3-t-butyl-5-methyl-4-hydroxyphenyl)propionamide], N,N'-bis[3-(3,5-di-t-butyl-4-hydroxyphenyl) droxybenzyl)isocyanurate, or 1,3,5-tris(4-t-butyl-3-hydroxy-2,6-dimethylbenzyl)isocyanuratel, and others. Among them, a phenolic compound having a t-butyl group (particularly, a plurality of t-butyl groups), in particular, a compound having a plurality of t-butylphenol sites, is preferred. These hindered phenol-series compounds may be used singly or in com-

bination.

[0045] The hindered amine-series compound may include a piperidine derivative having a steric hindrance group, for example, an ester group-containing pipendine derivative [for example, an aliphatic acyloxypiperidine (e.g., a Co. on aliphatic acyloxy-tetramethylpiperidine) such as 4-acetoxy-2,2,6,6-tetramethylpiperidine, 4-stearoyloxy-2,2,6,6-tetramethylpiperidine ndine or 4-acryloyloxy-2,2,6,6-tetramethylpiperidine; an aromatic acyloxypiperidine (e.g., a C7-11 aromatic acyloxy-tetramethylpiperidine) such as 4-benzoyloxy-2,2,6,6-tetramethylpiperidine; an aliphatic di-ortricarboxylic acid-bis-or trispiperidyl ester (e.g., a C_{2.20} aliphatic dicarboxylic acid-bispiperidyl ester) such as bis(2,2,6,6-tetramethyl-4-piperidyl) oxalate, bis(2,2,6,6-tetramethyl-4-piperidyl)malonate, bis(2,2,6,6-tetramethyl-4-piperidyl)adipate, bis(1,2,2,8,6-pentamethyl-4-piperidyl)adipate, bis(1,2,2,8,6-pentamethyl-4-piperidyl)adipate, bis(1,2,2,8,6-pentamethyl-4-piperidyl) ethyl-4-poper what indipate, bis(2,2,6,6-tetramethyl-4-piperidyl)sebacate, bis(1,2,2,6,6-pentamethyl-4-piperidyl)sebacate; an aromai o tetracarboxylic acid-bis- to tetrakispiperidyi ester (e.g., an aromatic di- or tricarboxylic acid-bis- or .er) such as bis(2,2,6,6-tetramethyl-4-piperidyl)terephthalate or tris(2,2,6,6-tetramethyl-4-piperidyl)benzene-1.3.5-tricarboxylate], an ether group-containing piperidine derivative [for example, a C1.10alkoxypiperidine (e.g., a C. alkoxy-tetramethylpiperidine) such as 4-methoxy-2,2,6,6-tetramethylpiperidine; a C_{6,8}cycloalkyloxy-piperidine such as 4-cyclohexyloxy-2,2,6,6-tetramethylpiperidine; an aryloxypiperidine such as 4-phenoxy-2,2,6,6-tetramethylpiperidine; a C₆₋₁₀aryl-C₁₋₄alkyloxy-piperidine such as 4-benzyloxy-2,2,6,6-tetramethylplperidine; or an alkylenedioxybisciperidine (e.g., a C₁₋₁₀alkylenedioxy-bispiperidine) such as 1,2-bis(2,2,6,6-tetramethyl-4-piperidyloxy)ethanel, an amide groupcontaining other ritine derivative [for example, a carbamoyloxypiperfoline such as 4-(phenyloarbamoyloxy)-2,2,6,6-tetramethylplperman a latkylenedioxy-bis piperidine substituted with a carbamoyloxy group, e.g., bis(2,2,6,6-tetramethyl-4piperidylines methylene-1,6-dicarbamate]. Moreover, the hindered amine-series compound may also include, for example, a polycondensate of piperidine derivatives having a high molecular weight (e.g., a polycondensate of dimethyl succinate and 1-(2-hydroxyethyl)-4-hydroxy-2,2,6,6-tetramethylpiperidine, and a poly(6-[(1,1,3,3-tetramethylpiperidine, and a poly(6-[(1,1,3,3,3-tetramethylpiperidine, and a poly(6-[(1,1,3,3,3-tetramethylpi no-1,3,5-triazin-2,4-d[y1][2-(2,2,6,6-tetramethylpiperldyl)amino] hexamethylene[4-(2,2,6,6-tetramethylpiperldyl)imino]]).These hindered amine-series compounds may be used singly or in combination.

[0046] These antioxidents may be used singly or in combination. The proportion of the antioxident may be about 0.001 to 5 parts by weight, preferably about 0.005 to 3 parts by weight, and more preferably about 0.01 to 2 parts by weight, relative to 100 parts by weight of the polyectal resin.

[0047] (Processing stabilizer)

The processing stabilizer may include at least one member selected from the group consisting of (a) a long-chain fatty add or a derivative thereof, (b) a polyoxyalkylene glycol, (c) a silicone compound, and others. [0048] (a) Long-chain or higher fatty add or derivative thereof

The long-chain or higher fatty acid may be a saturated fatty acid or an unsaturated fatty acid. Moreover, a part of hydrogen stoms in the higher fatty acid may be austirated with a substituent(s) such as hydroxyl group. Such a higher fatty acid may be exemplified by a mono- or di-fatty acid having not less than 10 carbon atoms, for exemple, a saturated of mono-fatty acid having not less than 10 carbon atoms, for exemple, a saturated of mono-fatty acid having not less than 10 carbon atoms for a saturated of the saturated of t

[0049] The derivetive of the higher fatty acid may include, for example, a fatty acid ester, a fatty acid areida, end others, 48 at the fatty acid ester, there is no particular imitation on its structure, and an ester of either a straight or branched chain fatty acid can be used. As the higher fatty acid ester, there may be mentioned, for example, an ester of the above-mentioned higher fatty acid with an alcohol (e.g., an ester having one or a plurality of ester bond(e), such as a moncester, a diester, a triester, or tetracterist. The alcohol constituting the higher fatty acid ester is not particularly initiated to applications. Such an alcohol may be a monohydric alcohol. As such an alcohol, a polyhydric alcohol is usually employed in an expresser.

[0050] The polyhydric alcohol may include a polyhydric alcohol having about 2 to 8 carbon atoms (preferably, about 2 to 8 carbon atoms) or a polymer thereof, for exemple, a diol exemptified by an akylene glycol [e.g., a Cz_eallylene glycol (preferably a Cz_eallylene glycol (preferably a Cz_eallylene glycol) such as ethylene glycol, dethylene glycol or propylene glycol) at riol exemplified by glycorin, ithrethylolpropene, or a derivative thereof; a tetraci exemplified by pentaenythitol, sorbitan, or a derivative thereof; as well as a homo- or copolymer of the polyhydric alcohol(e) [e.g., a homo- or copolymer of a polyoxyalkylene glycol such as a polyethylene glycol or a polypropylene glycol, a polygiverin, dipentiaerythritol, and a polypertaerythritol. The average degree of polymerization of the polyoxyalkylene glycol is not less than 2 (e.g., about 2 to 500), inclientally, in the

case of using the polyoxyalkylene glycol as a polyhydric alcohol, it is preferred to use, as a long-chain (or nigher) fatty acid constituting the eater, a fatty acid having not loss than 12 carbon atoms, for example, a saturated or unsaturated $C_{12\times 20}$ di-fatty acid. The abothols may be used singly or in combination.

- [0051] Examples of such an exter of a long-chain or higher fatty acid may include ethylene glycol mono- or dipamitate, ethylene glycol mono- or distantate, ethylene glycol mono- to triballinitate, glycenin mono- to triballinitate, glycenin mono- to trimantantate, pentate, pentate, genter glycol, ethylene glycol, ethylene glycol, ethically ethylene glycol (acut has a polyethylene glycol) ethylene glycol (acut has a polyethylene glycol), eth polyethylene glycol, eth glycol, eth glycol, eth glycol, ethylene glycol, ethyle
- [0052] Among these derivatives, as the fatty acid amide, for example, an acid smide (e.g., a monoamide and a 15 bisantile) of the liphor tatty acid is higher mono - of -farty acid) with an amine (such as a monoamine, a diamine or a polyamine) may be used. Among the acid amide, a bisamide is particularly preferred.
 - [0053] As the monoamide, there may be mentioned, for example, a primary acid amide of a saturated fatty acid (such as capit acid amide, lauric acid amide, myristic acid amide, palmitic acid amide, sitearic acid amide, erachic acid amide, behanic acid amide), a primary acid amide of an unsaturated fatty acid (such as oled amide); and a secondary acid amide of a saturated amide acid amide; and a secondary acid amide of a saturated and/or an unsaturated fatty acid with a monoamine (such as steary) stearic acid amide); and acid amide).
 - [0054] The bleamide may include, for example, a bleamide of the fatty acid with a C_{1-d}alkytenediamine (particulary, a C_{1-d}alkytenediamine). The concrete examples of the bleamide may include ethylenediamine-dipalmitic acid amide, a C_{1-d}alkytenediamine-diamide calcid amide, ethylenediamine-diamide calcid amide, ethylenediamine-diamide calcid amide, ethylenediamine-diamide, acid amide, and ethylenediamine-diamide. Evidenemine-diamide, ethylenediamine-diamide, propose are independently bonded to emine sites of an alkytenediamine, such as ethylenediamine, etiamic acid amide, and the such as ethylenediamine, etiamide, and exide to emine sites of an alkytenediamine, such as ethylenediamine, etiamide, and exide site acid amide, and the such as the such a
 - [0055] These long-chain (or higher) fatty acid amides or derivatives thereof may be used singly or in combination.
 [0056] (b) Polyoxyalkylene glycol
- The polyoxyalkylane glycol may include a homo- or copolymer of an alkylane glycol [e.g., a C_{2,e}alkylane glycol such as ethylane gycol, propylane glycol, or letramethylane glycol (preferably a C_{2,e}alkylane glycol), and a derivative (potor). [0057] Sepacific examples of the polyoyalkylane glycol and calculate a polyogo-gyankylane glycol), and a derivative glycol, a polyoropylane glycol or a polyearamethylane glycol (preferably a polyC_{2,e}acyalkylane glycol), a copolymer such as a polyoxyethylane-polyoxypropylane copolymer (e.g., a random or block copolymer), a polyoxyethylane-polyoxypropylane glyceryl ether, or a polyoxyethylane-polyoxypropylane monobutyl ether, and others. Among tham, the prefer on a includes a polymer having an oxyethylane-polyoxypropylane glycol), a polyoxyethylane-polyoxypropylane copolymer, and a derivative thereof.
- [0058] The number average molecular weight of the polyoxyalkylene glycol is about 3 x 10° to 1 x 10° (e.g., about 5 x 10°). The polyoxyalkylene glycols may be used singly or in combination.

 [0058] The number average molecular weight of the polyoxyalkylene glycols may be used singly or in combination.

 [0058] The number average molecular weight of the polyoxyalkylene glycols may be used singly or in combination.
 - The silicone-series compound may include a (poly)organosiloxane, and others. Examples of the (poly)organosiloxane may include, a monoorganosiloxane such as a dialkysiloxane (e.g., dimethysiloxane), an alkylarysiloxane (e.g., phe-fylimstylisioxane) or a darysiloxane (e.g., diphenysiloxane), a hormopolymer thereof (for example, a polymethysiloxane), or a copolymer thereof. Incidentally, the polyorganosiloxane may be an oll-
 - [0069] Moreover, the (poly)organosiloxane may include a modified (poly)organosiloxane (e.g., a modified silicone) having substituent(s) (such as an epoxy group, a nydroxyl group, an alkoxy group, a carboxyl group, an amining orgup or a substituted amining group (e.g., a disklyamining group), an either group, a vinyl group, or a (meth) acryloxyl group) in the end or main chain of the molecule. These silicone-series compounds may be used singly or in combination. [0061] The proportion of the processing stabilizer may be selected from, for example, about 0.001 to 10 parts by weight, preferably about 0.03 to 3 parts by weight, end more preferably about 0.03 to 3 parts by weight, relative to 100 parts by weight of the polyaceatir lessii. In particular, the proportion may be about 0.03 to 3 parts by weight.
 - [0062] (Heat stabilizer)
 The heat stabilizer may include (a) a basic nitrogen-containing compound, (b) a metal salt of an organic carboxylic acid, (c) an alkall or alkaline earth metal compound, (d) a hydrottalcite, (e) a zeolite, (f) a phosphine compound, and others, [0063] (a) Basic introcen-containing compounds and others.

As the basic nitrogen-containing compound (or basic nitrogen compound), at least one member selected from the group consisting of an aminotriazine compound, a guaridine compound, a urea compound, an amino acid compound, an amino alcohol compound, an imide compound, an amide compound, and a hydrazine compound may be used.

[0064] The aminotriazine compound may include melamine or a derivative thereof [e.g., melamine, and a condensate of melamine (melam, melam), guanamine or a derivative thereof, and an aminotriazine resin [for example, a co-polycondensation resin of melamine (e.g., a melamine-formativiga resin, a pheno-melamine resin, a melamine-phenotromatide hyde resin, a benzoguanamine-melamine rosin, and an aromatic polyamine-melamine resin), and a co-polycondensation resin of guanamine (e.g., a benzoguanamine-formatide hyde resin, and a benzoguanamine-phenot-formatide hyde resin).

[0065] Among the aminotriazine compounds, the derivative of guanamine may include an aliphatic guanamine compound [for example, a monoguanamine (e.g., a C1.24alkyl-substituted guanamine such as valeroguanamine, caproquanamine, heptanoguanamine, capryloguanamine, or stearoguanamine), and an alkylenebisguanamine (e.g., a C. o. alkylenebisguanamine such as succinoguanamine, glutaroguanamine, adipoguanamine, pimeloguanamine, suberoguanamine, azeloguanamine, or sebacoguanamine)], an alicyclic guanamine-series compound [for example, a monoguanamine (e.g., cyclohexanecarboguanamine, norbornenecarboguanamine, cyclohexenecarboguanamine, norbornenecarboguanamine, cyclohexenecarboguanamine, norbornenecarboguanamine, cyclohexenecarboguanamine, norbornenecarboguanamine, cyclohexenecarboguanamine, norbornenecarboguanamine, cyclohexenecarboguanamine, norbornenecarboguanamine, norbornecarboguanamine, norbornecarboguanamine, norbornecarboguanamine, norbornecarboguanamine, norbornecarboguanamine, n bornanecarboguanamine, and a compound obtained by introducing a functional group thereto (e.g., a derivative whose cycloalkane residue has one to three functional group(s) as a substituent, such as an alkyl group, a hydroxy group, an amino group, an acetoamino group, a nitryl group, a carboxyl group, an alkoxycarbonyl group, a carbamoyl group, an alkoxy group, a phenyl group, a cumyl group or a hydroxyphenyl group))], an aromatic guanamine-series compound (for example, a monoguanamine (e.g., benzoguanamine and a compound obtained by introducing a functional group thereto (e.g., a benzoguanamine derivative whose phenyl residue has one to five functional group(s) as a substituent, such as an alkyl group, an aryl group, a hydroxy group, an amino group, an acetoamino group, a nitryl group, a carboxyl group, an alkoxycarbonyl group, a carbamoyl group, an alkoxy group, a phenyl group, a cumyl group or a hydroxyphenyl group: for example, o-, m- or p-toluguanamine, o-, m- or p-xyloguanamine, o-, m- or p-phenylbenzoguanamine, o-, m- or phydroxybenzoguanamine, 4-(4'-hydroxyphenyl)benzoguanamine, o-, m-orp-nitrylbenzoguanamine, 3.5-dimethyl-4-hydroxybenzoguanamine, and 3,5-di-t-butyl-4-hydroxybenzoguanamine), α- or β-naphthoguanamine and a derivative obtained by introducing a functional group thereto, a polyguanamine (e.g., phthaloguanamine, Isophthaloguanamine, terephthaloguanamine, naphthalenediguanamine, and biphenylenediguanamine), and an aralkyl- or aralkyleneguanamine (e.g., phenylacetoguanamine, β-phenylpropioguanamine, and o-, m-orp-xylylenebisguanamine))], a hetero atomcontaining guanamine-series compound for example, an acetal group-containing guanamine (e.g., 2,4-diamino-6-(3,3dimethoxypropyl-s-triazine), a dioxane ring-containing guanamine (e.g., [2-(4',6'-diamino-s-triazin-2'-yl)ethyl]-1,3-dioxane, [2-(4',6'-diamino-s-triazin-2'-yl)ethyl]-4-ethyl-4-hydroxymethyl-1,3-dioxane}, a tetraoxospiro ring-containing guanamine (e.g., CTU-guanamine, and CMTU-guanamine), an isocyanuric ring-containing guanamine (e.g., 1,3,5-tris(2-(4', 6'-diamino-s-triazin-2'-yl)ethyljisocyanurate, and 1,3,5-tris[3-(4',6'-diamino-s-triazin-2'-yl)propyljisocyanurate), an imidazole ring-containing guanamine (e.g., guanamine compounds described in Japanese Patent Application Laid-Open Nos. 41120/1972 (JP-47-41120A)), and guanamine compounds described in Japanese Patent Application Laid-Open No. 154181/2000 (JP-2000-154181A)). Moreover, the aminotriazine compound may also include, for example, a compound which has alkoxymethyl group(s) on amino group(s) of the melamine, melamine derivative or quanamine-series compound [e.g., a mono- to hexamethoxymethylmelamine, a mono- to tetramethoxymethylbenzoguanamine, and a

[0066] The guanidine compound may include, for example, a non-cyclic guandine (e.g., glycocyamine, guanolin, guandine, and cyanoguandine), a cyclic guandine (e.g., glycocyamidine compound such as glycocyamidine, or creatinies; and cxalyiguandine or a cyclic guandine having a similar structure thereto, such as oxalyiguandine or cyclic diminographanic acidi; an inimi group-substituted urazole compound (e.g., iminourazole, and guanazole); an isocyanuric acid inide (e.g., isoammelide, and isoammelide, mallor/guandine, tratory/guandine; mesoxyliguandine; and these

[067] The urea compound may Indude, for example, a non-cyclic urea compound [for example, urea, an N-substituted urea having a substituent such as an alkyl group, a non-cyclic urea condersate (e.g., a polymer of urea, such as bitter, or bitures; and a condensate compound of urea and an aldehyde compound, such as methylenediurea or ureaform), a so cyclic urea compound [for example, a cyclic monourelide, e.g., a C-t_t-pailyleneurea (e.g., ettryleneurea, and crotonylideneurea), an aryleneurea (e.g., elmestin), a ureide of a fice-tokylic acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid for example, a hydratonic organical, and uramily, a ureide of a fi-aldehydic acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid for example, a hydratonic organical, and uramily, a ureide of a fi-aldehydic acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine, and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine), and urazole), a ureide of a charge of an c-hydroxy acid (e.g., uracil, thyrnine), and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine), and urazole), a ureide of an c-hydroxy acid (e.g., uracil, thyrnine), and urazole), a ureide of acid (e.g., uracil, thyrnine), and urazole), a ureide of acid (e.g., uracil, thyrnine), and urazole), a ureide of acid (e.g., uracil, thyrnine), and urazole), a ureide of acid (e.g., uracil, thyrnine urazole), a urazole), a ureide of a

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mono to octamethoxymethyl-CTU-quanaminel.

cantion such as 5,5-dibercy/lhydantolin; cC₁₋₁₀,alkylene-bishydantoln such as pentamethylenebishydantoin; and aliantoin or a metal as lithereof (cg., and a set such as aliantoin diflydioxysuluminum satt); a cycle diureite, for example, unic acid, an alkyl-substituted unic acid, acetyleneurea (glycolumi) or a derivative thereof (cg., a mono- to tetra (C₁₋₁ alkyl)glycolumi); conylidenedurea, a diureide of an or-hydroxy acid (cg., 1,1-methylenebis(5,5-dimethylthylaroth)), a diurea such as pruzatne, and a diureide of a dioxnoxylic acid (cg., a), alioxnntin, and purpuric acid).

yinyaeitomi), a durea such as p-urazine, and a durerie of a dearboxylic acid (e.g., alloxantin, and purpuric acids), ideas [[gosen]. The properties of the amino acid may include an e-amino acid (for example, a monaeminomocatboxylic acid (e.g., glycine, alanine, valin, norvalin, leucine, norleucine, isoteucine, priventylelanine, tyroshe, dilodotyrosine, surinamine, threenine, serine, proline, hydroxyprofine, hydrophan, methionine, cytine, eysteine, citalline, calminoturyica acid, ashabiyatropicolinic acid, teanine, and o- or m-tyrosine), a monoeminodicarboxylic acid (e.g., espartic acid, glutamic acid, asharydrodipicolinic acid, and hexatlyrocquinolinic acid), and a diaminomonocarboxylic acid (e.g., lysine, hydroxylysine, arginine, and hetidine)], a β-amino acid (e.g., β-amino bacid (e.g., β-amino acid (e.g., β-amino acid (e.g., β-amino acid (e.g., β-amino acid may also include an amino acid derivative in which a carboxyl group is subjected to metal salination (e.g., an alkali metal salt, an alkaline earth metal salt), and salt amino acid carboxyl con or esterification (e.g., methyl esterification, ethyl esterification, ethyl esterification).

[0069] The smino alcohol compound may include an amino C_{1,10}8liphatic mono- or polyol such as monoethenolamine, distributional mine, 2-amino-1-butanol, 2-amino-2-methyl-1-propanol, 2-amino-2-methyl-1,3-propanediol, 2-amino-2-ethyl-1,3-propanediol, or trieft/but/oxymethylaminomethane.

[0070] Examples of the imide compound may include an aromatic polycarboxylic acid imide such as phthalic acid imide, trimellitic acid imide, or pyromellitic acid imide, and others.

[071] The amide compound may include, for example, an alighatic carboxylic acid amide (e.g., malonamide, acipic acid amide, sebacic acid amide, and doscanedioic acid amide, so polyc carboxylic acid amide (e.g., e-caprolactam), an aromatic carboxylic acid amide (e.g., becaprolactam), any one state (e.g., accaprolactam), any one state (e.g., any one state (e.g.,

[0072] The hydrazine compound may include a carboxylic acid hydrazide different from the above-mentioned long-chain alliphatic carboxylic acid hydrazide, altoylic coachoxylic acid hydrazide, and dimer acid or trimer acid hydrazide. Examples of such a carboxylic acid hydrazide may include a short-chain aliphatic carboxylic acid hydrazide, ac C₂₊₁₈saturated or unsaturated fatty acid hydrazide such as adipic acid dihydrazide, lauric soid hydrazide, sebacic acid dihydrazide, cobecensicion acid dihydrazide, or sorbic acid hydrazide profession policy acid hydrazide, per acid hydrazide, acid hydrazid

The metal sat of the organic carboxylic acid may include, for example, a self of an organic carboxylic acid with a metal (e.g., an alkali metal such as U.N ac rk; an alkaline earth metal such as Mg or Ca; and a transition metal such as Zn).

[9074] The organic carboxylic acid may be a compound of low molecular weight. As the organic carboxylic acid, there may be used a saturated or unsaturated lower aliphatic carboxylic acid, and the example of the molecular weight. As the organic carboxylic acid, and action, and acid, there may be used a saturated or unsaturated lower aliphatic carboxylic acid, and acid, acid,

[0075] The unsaturated lower aliphatic carboxylic acid may include, for example, an unsaturated C₃₋₄monoccarboxylic acid (e.g., (meth)acrylic acid, crotonic acid, and isocrotonic acid), an unsaturated C₄₋₄dicarboxylic acid (e.g., maleic acid, and fumaric acid), and a hydroxy acid thereof (e.g., propoid) acid).

[0076] Moreover, exemplified as the polymer of the unsaturated aliphatic carboxylic acid may be a copolymer of a

polymerizable unsaturated carboxylic acid [for example, an $\alpha_i\beta$ -ethylene-type (ethylenic) unsaturated carboxylic acid, for example, a polymerizable unsaturated monocarboxylic acid (such as (meth)pacrylic acid), a polymerizable unsaturated polyvaroxylic acid (such as fiscancia acid, mainted acid, an acid antyvinde of the polyearboxylic acid, or a monocaster of the polyearboxylic acid (e.g., a monoC₁₋₁,alkyl ester of the polyearboxylic acid (e.g., a monoC₁₋₁,alkyl ester of the polyearboxylic acid such as monoethyl mailastal) with an olefin (e.g., an CC_{2-1} ,polefin such as ethylene or provivene).

[0077] These metal saits of the organic carboxylic acids may be used singly or in combination,

[0078] The preferred metal salt of the organic carboxytic acid may include a salt of an organic carboxytic acid with an alkali metal (e.g., lithium citrate, potassium citrate, sociaum citrate, it into many carbox, and thinking 12-hydroxystearate), a salt of an organic carboxytic acid with an alkaline earth metal (e.g., magnesium aceate, calcium aceate, magnesium citrate, calcium citrate, calcium citrate, calcium stearate, magnesium stearate, magnesium 12-hydroxystearate, and calcium 12-hydroxystearate), and nonomer resin ic resin in which at least a part of carboxyt groups contained in the copolymer of the polymer/salterate), and nonomer resin ic resin in which at least a part of carboxyt groups contained in the copolymer of the polymer/salterate), and calcium contained in the copolymer of the polymer/salterate polycarboxytic acid with the olefin is neutralized with an in on of the motal, and others. The looner resin is, for example, commercially available as ACLYN (manufactured by Du Pont, and others.)

[0079] Among the metal salts, in view of stabilizing effects, a salt with an alkaline earth metal such as calcium citrate, magnesium stearlet, calcium stearate, magnesium 12-hydroxystearate, or calcium 12-hydroxystearate (particularly, calcium citrate) is preferen.

[0080] (c) Alkalin c alkaline earth metal compound The alkali or alkaline earth metal compound may include an inorganic compound exemplified by a metal oxide (such as CaO, or MgO), a metal hydroxide (such as LOH, Ca(CH)₂, or Mg over the control of the control of the control oxide of the control oxide ox

25 As the hydrotalicite, hydrotalicites recited in Japanese Patent Application Laid-Open No. 1241/1985 (JP-80-1241A) and Japanese Patent Application Laid-Open No. 59475/1997 (JP-9-59476A), such as hydrotalicite compounds represented by the following formula are usable.

[0083] $[M^{2+}_{1-x}M^{3+}_{x}(OH)_{2}]^{x+}[A^{n-}_{x/n}\cdot mH_{2}O]^{x-}$

In the formula, M²⁺ represents Mg²⁺, Mn²⁺, Fe²⁺, Co²⁺, or any of other divalent metal ions; M³⁺ represents Al²⁺, Fe²⁺, or any of other invalent metal ions; A represents CO₂²⁺, OH, HPO₄²⁺, SO₄²⁺, or any of other n-valent anions (particularly, monovalent or divalent anion); x is O< x < 0.5; and m is 0.5 m < 1.

These hydrotalcites may be used sindly or in combination.

[0084] incidentally, the hydrotalcite is available from Kyowa Chemical Industry Co., Ltd. under the trade name "DHT-4A-2", or "Alcamizer".

[0085] (e) Zeolite

The zeolite is not particularly limited to a specific one, and a zeolite other than H-type zeolite can be employed, for exampe, a zeolite recited in Japanese Patent Application Laid-Open No. 62 142/1955 [JP-7-62-142A] | zeolites the small-set unit cell of which is a crystalline aluminosilicate with an alkaline and/or alkaline earth metal (A, X, Y, L, and ZSM-type zeolites, mordenite-type zeolites, mordenite-type zeolites, mordenite-type zeolites).

[0086] These zeolites may be used singly or in combination.

[0087] Incidentally, A-type zeolite is available as "ZEOLAM-series (A-3, A-4, A-5)", "ZEOSTAR-series (KA-100P, NA-100P, CA-100P)" or others, X-type zeolite as "ZEOLAM-series (F-9)", "ZEOSTAR-series (NX-100P)" or others, and Ytype zeolite as "HSZ-series (320NAA)" or others, from Tosoh Corp. or Nippon Chemical Industrial Co., Ltd. [0088] (f) Phosphine compound

Examples of the phosphine compound may include a phosphine compound such as an allylphosphine (for example, a rtC_{1-toalkiy}(phosphine such as triethyphosphine, tripropylphosphine, or tributyphosphine), a cycloakiyphosphine (for example, a triC_{2-toa}(phosphine which may have a substituent (such as en amino group or a C_{1-ta}(ky) group), such as triphenylphosphine, p-tolyldiphenylphosphine, di-p-tolylphosphine, tri-m-aminophenylphosphine, int(2,4-dimethyphosphine), an artilyphosphine (for example, as

a tri(C₆₊₂eny(C₁₋₄elky))phosphine such as tri(C₆₊₂eny(C₁₋₄elky))phosphine), an equippinosphine (or example, a tri(C₆₊₂eny(C₁₋₄elky))phosphine such as participation (or example, a monoor of C₆₊₂eny(C₁₋₄elky))phosphine (or example, a monoor of C₆₊₂eny(C₁₋₄elky))phosphine (or example, a mono-or of C₆₋₂eny(C₁₋₄elky))phosphine (or example, a mono-or of C₆₋₂eny(C₆₋₂eny(C₆₋₂eny(C₆₋₂eny))phosphine (or example, a mono-or of C₆₋₂eny(C₆₋₂eny(C₆₋₂eny))phosphine (or example, a mono-or of C₆₋₂eny(C₆₋₂eny))phosphine (or example, a mono-or of C₆₋₂eny)phosphine) or example, and a C₆₋₂eny(C₆₋₂eny(C₆₋₂eny)phosphine) or example, a bleigh(C₆₋₂eny)phosphine) or example, a bleigh(C₆₋₂eny)phosphine) or in combination (or in combination).

[0089] These heat stabilizers may be used singly or in combination. In particular, in the case of using the basic nitrogen-

containing compound in combination with at least one member selected from the group consisting of the metal salt of an organic carboxylic acid, the aixfall or alkaline earth metal compound, the hydrotaliste, the zeolite, and the phasphine compound, heat stability can be also impaired to the resin composition at a smaller amount of the heat stabilizer, inclidentally, the resin composition of the present invention can also improve heat stability without substantially containing a phosphorus containing fairar partagrant.

[0900] In the case where the resin composition contains the heat stabilizer, the proportion of the heat stabilizer may be, for example, selected from the range of about 0.001 to 10 parts by weight, and preferably about 0.001 to 5 parts by weight (perficularly about 0.01 to 2 parts by weight, leafable to 100 parts by weight of the polyscated resin. Incidentally, among the heat stabilizers, the hydrazine compound (the short-chial aligneds carboxylic acid hydrazine) and reperately used at a small amount because there is a possibility that to large amount of such a hydrazine compound brings about bleeding out thereof from the polyscatal resin composition or deterioration in processing stability of the resin composition. The proportion of the hydrazine compound may be usually not more than 1 part by weight (e.g., about 0.01 to 1 part by weight), preferably about 0.05 to 0.8 parts by weight of the polyscated resin.

[0091] (Weather (light)-resistant stabilizer of the weather (light)-resistant stabilizer may include (a) a benzotriazole-series compound, (b) a benzotrenene-series compound, (c) an aromalic benzotre-series compound, (d) a cyanoacyjste-series compound, (e) an oxalic anlide-series compound, (f) a hydroxyaryl-1,5,5-riazine-series compound, (g) a hindered amine-series compound, and others. [0092] (a) Benzorriazole-series compound

20 Examples of the benzotriazole-senies compound may include a benzotriazole compound having an any group substituted with a hydroxyl group and a C-4_ellivig group, such as 2-(2'-hydroxy-5'-methytchenyl)benzotriazole, 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole or 2-(2'-hydroxy-3', 5'-di-ellivenyl)benzotriazole or 2-(2'-hydroxy-3', 5'-di-ellivenyl)benzotriazole or 2-(2'-hydroxy-3', 5'-di-ellivenyl)benzotriazole or 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole or 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole or 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole or 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole (a benzotriazole or 2-(2'-hydroxy-3', 5'-dil-enyl)benzotriazole (a benzotriazole or 2-(2'-hydroxy-4'-cotoxyphenyl)benzotriazole or 2-(2'-hydroxy-4'-cotoxyphenyl)benzotriazole and others.

[0093] Among these benzotriazole-series compounds, the particularly preferred one includes a benzotriazole compound having a C₆₋₁₀87l (particularly, phenyl) group substituted with a hydroxyl group and a C₆₋₂₀8lyl group, as well as a benzotriazole compound having an anyl group substituted with a hydroxyl group and a C₆₋₁₀8ryl-C₁₋₅8ikyl (particularly, phenyl-C₁₋₅8ikyl group.

(0094) (b) Benzophenone-series compound

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Exemplified as the benzophenone-series compound may be a benzophenone compound having a plurality of hydroxy groups (e.g., a di-to tetrahydroxybenzophenone such as 2,4-dihydroxybenzophenone; a benzophenone compound having a hydroxyl group, and an anyl or aralkyl group substituted with a hydroxyl group, such as 2-hydroxy4-arboxyzylidanzophenone); a benzophenone compound having a hydroxyl group and an alkoxyl (C_{1+t}alkoxy) group (e.g., 2hydroxy4-drethoxybenzophenone, 2-hydroxy4-doctoxybenzophenone, 2-hydroxy4-doctoxybenzophenone, 2-hydroxy4-doctoxybenzophenone, 2-hydroxy4-doctoxybenzophenone, 2-hydroxy4-doctoxybenzophenone, 2-hydroxy4-methoxy6-sulfobenzophenone); and others.

[0095] Among these benzophenone-series compounds, the benzophenone-series compound preferably includes a 40 benzophenone compound having a hydroxyl group, and a C₆₋₁₉agyl (or C₆₋₁₉agyl-C₁₋₄alikyl) group substituted with a hydroxyl group, and a phenyl-C₁₋₄alikyl group substituted with a hydroxyl group, [0096] (c) Aromatic benzoate-series compound

The aromatic benzoate-series compound may include an alkylarytsalicylate such as p-t-butylphenylsalicylate of p-octylphenylsalicylate (particularly, an alkylphenylsalicylate).

[0097] (d) Cyanoacrylate-series compound

Exemplified as the cyanoacrylate-series compound may be a cyano group-containing diarylacrylate such as 2-ethylhexyl-2-cyano-3,3-dipherylacrylate or ethyl-2-cyano-3,3-dipherylacrylate (particularly, a cyano group-containing dipherylacrylate).

[0098] (e) Oxalic anilide-series compound

The oxalic anilide-series compound may include, for example, an oxalic dismide compound having an any group (such as phenyl group) on a nitrogen atom in which the any group may have a substituent(s), exemplified by N-(2-ethylphenyl)-N-(2-ethoxy-p

Examples of the hydroxyan/i-1,3.5-triazine-series compound may include a 2.4-diC₈₋₁₄₈n/i-6-(mono- or dihydroxyC₆₋₁₄an/i-1,3.5-triazine life example, a 2,4-diC₆₋₁₀₈n/i-6-(mono- or dihydroxyC₆₋₁₄₈n/i)-1,3.5-triazine which may have a substituentia) (such as a C₁₋₁₀₈liky) group, a C₁₋₁₆likoy group, a C₁₋₁₆likoy group, or a C₆₋₁₀₈n/iC₆, alloxy group) on an any group thereof, e.g., a hydroxyan/titazine such as 2.4-diphen/i-6(2-4-dihydroxyhen/i-1,3.5-triazine); a hydroxyan/titazine such as 2.4-diphen/i-6(2-4-dihydroxyhen/i-6(2-4-dihydroxyhen/i-1,3.5-triazine); a hydroxyan/titazine such as 2.4-diphen/i-6(2-4-dihydroxyhen/i-1,3.5-triazine); a hydroxyan/titazine such as 2.4-diphen/i-6(2-4-dihydroxyhen

alkoxyaryltriazine such as 2,4-diphenyl-6-(2-hydroxy-4-methoxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4ethoxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4-propoxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4-propoxyphenyl-6-(2-hydr butoxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4-hexyloxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4octyloxyphenyl)-1,3,5-triazine, 2,4-diphenyl-6-(2-hydroxy-4-dodecyloxyphenyl)-1,3,5-triazine, or a 2,4-di(p-tolyl or 2',4'dimethylphenyl)-6-(2-hydroxy-C₁₋₁₆alkoxyphenyl)-1,3, 5-triazine corresponding to each of these 2,4-diphenyl-6-(2-hydroxy-alkoxyphenyl)-1,3,5-triazines; a hydroxyaralkyloxyaryltnazine such as 2,4-diphenyl-6-(2-hydroxy-4-benzyloxyphenyl)-1,3,5-triazine, or 2,4-di(p-tolyl or 2,4'-dimethylphenyl)-6-(2-hydroxy-4-benzyloxyphenyl)-1,3,5-triazine; a hydroxyalkoxyalkoxyaryltriazine such as 2,4-diphenyl-6-(2-hydroxy-4-(2-butoxyethoxy)phenyl)-1,3,5-triazine, or 2,4-di-ptolyl-6-(2-hydroxy-4-(2-hexyloxyethoxy)phenyl)-1,3,5-triazine; and others. Among these compounds, the hydroxyphenyl-1,3,5-triazine-series compound is preferred.

[0100] (g) Hindered amine-series compound

As the hindered amine-series compound, the hindered amine-series compound as exemplified in the paragraph of the above-mentioned antioxidant may be used,

[0101] These weather (light)-resistant stabilizers may be used singly or in combination. The same or different species of the weather (light)-resistant stabilizers may be used in combination.

[0102] Incidentally, it is preferred to use the hindered amine-series compound (g) in combination with other weather (light)-resistant stabilizer. In particular, it is preferred to use the benzotriazole-series compound (a) in combination with the hindered amine-series compound (g). The proportion (weight ratio) of the hindered amine-series compound relative to other weather (light)-resistant stabilizer(s) (particularly, the benzotrlazole-series compound) [the hindered amineseries compound (g)/other weather (light)-resistant stabilizer(s)] may be, for example, about 0/100 to 80/20, preferably about 10/90 to 70/30, and more preferably about 20/80 to 60/40.

[0103] The proportion of the weather (light)-resistant stabilizer is, for example, about 0 to 5 parts by weight (e.g., about 0.01 to 5 parts by weight), preferably about 0.1 to 4 parts by weight, and more preferably about 0.1 to 2 parts by weight. relative to 100 parts by weight of the polyacetal resin.

[0104] (Coloring agent)

As the coloring agent, various dyes or pigments may be used. As the dye, a solvent dye is preferred, and includes, for example, an azo-series dye, an anthraquinone-series dye, a phthalocyanine-series dye or a naphthoguinone-series dye. The pigment may be an inorganic pigment or an organic pigment.

[0105] Exemplified as the inorganic pigment may be a titanium-senes (titanium-containing) pigment, a zinc-series (zinc-containing) pigment, a carbon black (e.g., a furnace black, a channel black, an acetylene black, and Ketjen black), an iron-series (iron-containing) pigment, a molybdenum-series (molybdenum-containing) pigment, a cadmium-series (cadmium-containing) pigment, a lead-series (lead-containing) pigment, a cobalt-series (cobalt-containing) pigment, and an aluminum-series (aluminum-containing) pigment.

[0106] The organic pigment may include an azo-series pigment, an anthraquinone-series pigment, a phthalocyanineseries pigment, a quinacridone-series pigment, a perylene-series pigment, a perinone-series pigment, an isoindolineseries pigment, a dioxazine-series pigment, or a threne-series pigment.

[0107] The coloring agent may be used singly, or a plurality of these coloring agents may be used in combination. The use of a coloring agent having a high fight-shielding effect [such as a carbon black, a titanium white (a titanium oxide), a phthalocyanine-series pigment, a perylene-series pigment (particularly a carbon black, a perylene-series black pigment)] ensures improvement in weather (light)-resistance of the polyacetal resin composition.

[0108] The content of the coloring agent is, for example, about 0 to 5 parts by weight (e.g., about 0.01 to 5 parts by weight), preferably about 0.1 to 4 parts by weight, and more preferably about 0.1 to 2 parts by weight, relative to 100 parts by weight of the polyacetal resin.

[0109] To the polyacetal resin composition of the present invention may be optionally added a conventional additive (s) singly or in combination. Examples of the additive may include an antioxidant (e.g., a phosphorus-containing, a sulfurcontaining, a hydroquinone-senes, and a quinoline-senes antioxidant), a specific carboxylic acid (e.g., a carboxylic acid described in Japanese Patent Application Laid-Open No. 239484/2000 (JP-2000-239484A)), an impact resistance improver [e.g., an acrylic core-shell polymer, a polyurethane-series resin, and a polyester-series resin], a slip-improving agent [e.g., an olefinic polymer, a silicone-series resin, and a fluorine-containing resin], a mold-release agent releasing agent), a nucleating agent, an antistatic agent, a flame retardant, a foaming agent, a surfactant, an antiba agent an antifungal agent, an aromatic agent, a perfume, various polymers [e.g., an across resin (a homo-or co her of s. C_{1.40}alkyl (meth)acrylate such as a poly(methyl methacrylate)), a polycarbonate-series resin, a polyclef

or resin, a polyvinyl alcohol-series resin, and an aliphatic polyester-series resin (e.g., a poly(L-lactic acid), a 201, (D-lactic acid), a poly(D/L-lactic acid), a polyglycolic acid, and a copolymer of glycolic acid and lactic acid (e.g., D-, L-or D/L-lactic acid))], a filler, and others.

[0110] Moreover, if necessary, the resin composition may be further blended with one or combination of a conventional filler (such as a fibrous, plate-like or particulate filler) to improve properties of the shaped article of the present invention. Examples of the fibrous filler may include an inorganic fiber (e.g., a glass fiber, a carbon fiber, a boron fiber, and a

potassium titanate fiber (whisker)), an organic fiber (e.g., an amide fiber), and others. As the plate-like filler, there may be mentioned a glass flake, a mica, a graphite, a variety of metal foll, and others. Examples of the particulate filler may include a metal oxide (e.g., zinc oxide, and alumina), a sulfate (e.g., calcium sulfate, and magnesium sulfate), a carbonate (e.c., calcium carbonate), a glass (e.g., a milled fiber, a glass bead, and a glass balloon), a silicate (e.g., a taic, a kaolin, a silica, a diatomite, a clay, and a wollastonite), a sulfide (e.g., molybdenum disulfide, and tungsten disulfide), a carbide (e.g., graphite fluoride, and sillcon carbide), boron nitride, and others.

[0111] (Production process of polyacetal resin composition)

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The polyacetal resin composition of the present invention may be a particulate mixture or a molten mixture, and it can be prepared by mixing a polyacetal resin with the specific carboxylic acid hydrazide (the long-chain aliphatic carboxylic acid hydrazide and/or the alloyofic carboxylic acid hydrazide), and if necessary, other additive(s) [e.g., a stabilizer (an antioxidant, a processing stabilizer, a heat stabilizer, a weather (light)-resistant stabilizer), an impact resistance improver, a slip-improving agent, a coloring agent and/or a filler], in a conventional manner. For example, (1) a process comprising feeding all components through a main feed port, kneeding and extruding the resulting mixture into pellets with an extruder (e.g., a uniaxial or biaxial extruder), and molding a shaped article from the pellets, (2) a process comprising feeding component(s) (e.g., a polyacetal resin, and the above-mentioned other additive(s)) free from the specific carboxylic acid hydrazide through a main feed port, feeding component(s) containing at least the specific carboxylic acid hydrazide (as other component(s), there may be mentioned a polyacetal resin, the above-mentioned other additive(s), or the like) through a side feed port, kneeding and extruding the resulting mixture into pellets with an extruder, and molding a shaped article from the pellets, (3) a process comprising feeding component(s) containing a part of the specific carboxylic acid hydrazide (as other component(s), a polyacetal resin, other additive(s), or the like) through a main feed port and feeding component(s) containing the residual specific carboxylic acid hydrazide, and, if necessary, a shortchain aliphatic carboxylic acid hydrazide and/or an aromatic carboxylic acid hydrazide as a heat stabilizer (as other component(s), a polyacetal resin, other additive(s), or others) through a side feed port, kneading and extruding the fed components by using an extruder to prepare pellets, and molding a shaped article from the pellets; (4) once making pellets (master batch) different in formulation, mixing (diluting) the pellets in a certain ratio, and molding a shaped article having a certain formulation from the resulting pellets, or (5) a process comprising allowing the specific carboxylic acid hydrazide to coexist with or adhere to a pelletized polyacetal resin by for example spraying or coating (e.g., surfacecoating), and molding a shaped article having a certain formulation from the resulting pellets is utilized.

[0112] Among these processes, the processes (1), (2) and (3) are preferred. In particular, it is preferred to melt-knead components by a uniaxial or biaxial extruder having exhaust (or degas) vent port(s) of not less than 1. Moreover, the carboxylic acid hydrazide may be side-fed through either of a feed port of the upstream or downstream of an exhaust vent port. In particular, it is preferred to feed the carboxylic acid hydrazide from the downstream of the exhaust vent port. Further, in the extruding and preparing step, the amount of formaldehyde emitted from the obtained shaped (or molded) article can be further reduced by a preparation method comprising preblending a processing auxiliary such as water and/or an alcohol (e.g., methanol, ethanol, isopropyl alcohol, and n-propyl alcohol) or infusing the processing auxiliary through a feed port of the upstream of an exhaust vent port, and exhausting and removing volatile component (s) containing water and/or the alcohol from the exhaust vent port. The amount of water and/or the alcohol to be added as such a processing auxiliary is not particularly limited to a specific one. The amount of water and/or the alcohol may be usually selected from the range of about 0 to 20 parts by weight relative to 100 parts by weight of the polyacetal resin. and may be preferably about 0.01 to 10 parts by weight and more preferably 0.1 to 5 parts by weight relative to 100

parts by weight of the polyacetal resin.

[0113] Incidentally, in the preparation of a composition for use in a shaped article, mixing of a powdered (particulate) polyacetal resin as a substrate (e.g., a powder (particulate) obtained by grinding a part or all of the polyacetal resin) with other components (e.g., the specific carboxylic acid hydrazide, other additive(s) (e.g., a stabilizer, an impact resistance improver, a slip-improving agent, a coloring agent and/or a filler) followed with melt-kneading improves the degree of dispersion of the additives and therefore is advantageous.

[0114] The polyacetal resin composition of the present invention realizes that the emission of formaldehyde due to oxidation or thermal decomposition or the like of the polyacetal resin is remarkably restrained or inhibited and that the working environment is improved or ameliorated particularly in the molding and processing (particularly, a melt-molding and processing) step. Moreover, deposition of decomposition products or additives on the mold (mold deposit), blooming or bleeding of such products or additives from a shaped article can be remarkably restricted or inhibited, and various problems on the molding and processing step can be overcome.

[0115] (Shaped article)

The present invention also includes a shaped article formed from the resin composition. The shaped article contains the polyacetal resin and the specific carboxylic acid hydrazide in combination, and has excellent stability in an extrusion and/or molding process with having extremely small amount of emission (or generation) of formal dehyde. In other words, shaped articles molded from the conventional polyacetal resins containing antioxidants and other stabilizers liberate relatively large amounts of formaldehyde, cause corrosion and discoloration, as well as pollute the living and working

environment. For example, the formatidelyde emission from commercial ordinary polyacetal resin articles is about 2 to 5 µg per one cm²0 startice area under dry conditions (in a constant-temperature dry atmosphere) and/or about 3 to 6 µg per one cm²0 of surface area under humis conditions (in a constant-temperature moisture-laden atmosphere)

[0]18] On the other hand, in the polyacetal resin shaped article of the present invention, the amount of formaldehyde of emission from the shaped article can be effectively reduced by a smaller amount of the specific carboxylic acid hydrazide and the heat stabilizer (formaldehyde inhibitor) in combination, the amount of formaldehyde emission can be also inhibited to a large extent. Concretely, the amount of the formaldehyde emission is not more than 1.5 gip ero nor cm² of surface area of the shaped article under dry conditions, preferably about 0 to 1.0 µg, more preferably about 0 to 0.6 µg, and usually about 0.001 to 1.0 µg, and further, about 0 to 0.1 µg is also achievable. Moreover, in humid conditions, the formaldehyde emission is not more than 2.5 µg (e.g., about 0 to 2.2 µg) per one cm² of surface area of the shaped article, preferably about 0 to 1.2 µg, more preferably about 0 to 0.4 µg, and further, about 0 to 0.2 µg is also achievable. The amount in humid conditions may be usually about 0.001 to 1.2 µg.

[0117] The shaped article of the present invention may show the above-mentioned formaldehyde emission under either dry conditions or humid conditions. In particular, the shaped article shows the above formaldehyde emission level under both dry and humid conditions in many cases. Therefore, the shaped article of the present invention can be used as a material which can be adapted to more severe environment.

[0118] The formaldehyde emission under dry conditions can be determined as follows.

[0119] After the shaped article of polyacetal resin is out if necessary and its surface area is measured, a suitable portion of the article (e.g., the amount equivalent to a surface area of about 10 to 50 cm²) is placed in a vessel (20 mL capacity) to seal and stand for maintained at a temperature of 80°C for 24 hours. Then, this sealed vassel is charged with 5 mL of water and the formaldehyde in the aqueous solution is assayed in accordance with JIS (dapanese Industrial Standards) K0102, 28 (under the heading of Formaldehyde) to calculate the formaldehyde emission per unit surface area of the shaped article (µg/m²n²).

25 [0120] The formaldehyde emission under humid conditions can be determined as follows.

[011] After the shaped article of a polyacetal resin is cut if necessary and its surface area is measured, a suitable portion of the shaped article (e.g., the amount equivalent to a surface area of about 10 to 100 cm²) is suspended from the ld of a sealable vessel (1 L capacity) containing 50mL of distilled water. After seal of the vessel, the vessel is allowed to stand (or maintained) in a constant temperature oven at 50°C for 3 hours. Thereafter, the vessel is allowed to stand at a room temperature for 1 hour and the formatidehyde in the aquecus solution in the vessel is assayed in accordance with US K0102, 29 (under the heading of Formaldehyde) to celculate the formaldehyde amission per unit surface area of the article (µg/cm²).

[0122] The above quantitative definition on formaldehyde emission in the present invention is adeptable as far as the powdered at resin and the specific actoxylic acid hydrazide, that is, the definition is adeptable not only for shaped anticles molded from polyaciest resin compositions compositions compositions composition and a mold-release agent), but also for sheped articles molded from comparable resin compositions containing an inorganic filler and/or other polymens, even if only a major part of the surface of the article (for example, 50 to 100% of the total surface area) is constituted by the polyacetal resin (for example, a multi-colored article) or contact article).

40 INDUSTRIAL APPLICABILITY

[0123] The resin composition of the present invention is useful for molding various shaped articles by a conventional molding (or shaping) method (for example, hijection molding, extrusion molding, compression molding, blow molding, vacuum molding, froat molding, creation molding, and gas injection molding).

45 [0124] Moreover, the shaped article (or moded article) of the present invention finds application in any field of use where formatidehyde is objectionable (e.g., inch and inlever as bloycle parts) and can also be used advantageously as parts and members in a variety of fields inclusive of automotive parts, electrical and electronic component (integrate) and one members and pipeline installation parts, household (for daily use) and comende product parts, and medical device (for diagnostic or therapeutic use) parts.

[0125] More specifically, the automotive parts may include car interior parts such as inner handle, fuel trunk opener, seat bet buckle, assisting, verillers, whiches, knob, lever, and clip; electrical system parts such as meters and connectors; in-vehicle electrical and electrical parts in contract with metals, typicall; in evindow regulator certifier parts, in contract with metals, typicall; in evindow regulator certifier plate, mechanical parts such as door lock actuator parts, mirror parts, where motor system parts, and turb system parts.

[0126] The electrical or electronic component parts (the mechanical parts) may include, for example, parts or members constituted with shaped articles of polyacetal resin and fitted with a number of metal contacts [e. g. audio equipment such as oxideo tape recorder (VTR), 8 mm or or other video camera, etc., office automation (OA) equipment such as copying machines, facsimile, word processor, computer, toys actuating the component of the content of the

the driving force of an electric motor or a spring, a telephone, a keyboard as an accessory to a computer or the like]. To be specific, there can be mentioned chassis (base), gear, lever, cam, pulley, and bearing. Furthermore, the electrical or electrical component parts are applicable to optical and magnetic recording medium parts at least party made or moided polyacetal resin (e.g. metal trihr-film magnetic tape cassette, magnetic disk carridge, opticomagnetic disk carridge, acci, jain more particularly, the metal tape cassette for music, digital audio tape cassette, in mm video tape cassette, floppy (registered trademark) disk carridge, entindisk carridge, etc.) as specific optical and magnetic medium parts, there can be mentioned tape cassette parts (tape cassette body, reel, hub, guide, roller, stopper, lid, etc.) and disk carridge parts (disk carridge body (cass), whater, cramping plate, etc.).

[0127] In addition, the shaped article of a polyacetal resin according to the present invention can be used with advantage in architectural members and pielinic parts such as lighting equipment parts, interior architectural members (such as fittings, fixtures, furnishings), piping, cock, facute, rest from (watorby-related parts, etc., a broad range of products related to daily living, cosmetic products, and medical devices, for example factoner (such as elide fastener, snap fastener, hoop-and-loop fastener, rail fastener), stationery, chepstick or injectic cases, cleansing device, water cleaner, spray nozyce, syray device or container, sensot container, general vessels, and syringe holder.

EXAMPLES

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[0128] The following examples are intended to describe this invention in further detail and should by no means be interpreted as defining the scope of the invention,

20 [0129] Incidentally, referring to the examples and comparative examples, the moldability (the amount of the deposit on the mold), the amount of formaldehyde emission from the molded (or shaped) articles under dry and humid conditions, and the bleeding property were evaluated based on the following methods. [0130] [Moldability (the amount of the deposit on the mold)]

A pellet formed with a polyacetal resin composition was continuously or successively shaped or molded by using a 30 5 th injection molding machine (100 shots) to cotain a certain-shaped article (20 mm in diameter and 1 mm in thickness), and the degree of the deposition on the mold was evaluated and classified into five grades, incidentally, the larger the number of the levels is, the lower or smaller the amount of the deposit (i.e., mold deposit) is.

[0131] [Amount of formaldehyde emission from shaped article in dry conditions]

Each resin sample consisting of 10 test pleces (one test plecer 2 mm x 2 mm x 50 mm; total surface area: about 40 cm²) 30 was pleced in a vessel (capacity 20 mL) to seal and heated in a constant temperature over at 80°C for 24 hours. After air-cooling to a room temperature, 5 mL of distilled water was injected into the vessel using a syringe. The formaldehyde content of this acueous solution was ostermined in accordance with JIS KO102, 29 (under the heading of Formaldehyde) and the formaldehyde gas emission per surface area (µg/cm²) was calculated.

[0132] [Amount of formaldehyde emission from shaped article in humid conditions, and the bleeding property]

Two plate test pleces (one plece: 100 mm x 40 mm x 2 mm; total surface area of 8.5 6 m²) were suspended from a lid of a polyethylene bottle (capacity 1 L) containing 50 mL of distilled water. The bottle was sealed to stand in a constant temperature oven at 80°C for 3 hours, followed by standing for 1 hour at room temperature. The formalidehyde content of the aqueous solution in the bottle was determined in accordance with JIS K0102, 29 (under the heading of Formalidehyde) and the formalidehyde gas emission per surface area of the article (agrand) was calculated.

0 [0133] Further, the surface of the plate test piece (shaped article) after the test was visually observed, and the degree of the bleeding was evaluated based on the following orderta.

- "A": No bleeding was observed.
- "B": Slight bleeding was observed.
- "C": Extremely heavy bleeding was observed.

[0134] Examples 1 to 14 and 16 to 19

100 parts by weight of a polyscetal resin copolymer, a carboxylic soid hydrazide, an entioxidant, a processing stabilizer, a heat stabilizer, a coloring agent, and a weather (fight)-resistant stabilizer in the proportions indicated in Tables 1 and 2 were problemed (or premixed). Concerning each of thus obtained mixtures, the mixture was supplied moyel a main feed port of a binxial extruder (50 mm diameter) having one vent port, and met-mixed to prepare a pelletized composition. From thus obtained pellets, prescribed test pieces were fabricated with an injection modifing machine, and concerning each test piece, the amount of formalder/yde emission from the test piece, and the bleeding property were measured. The results are shown in Tables of and 2.

55 [0135] Example 15

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Es parts by weight of a polyacetal resin copplymer, an antioxidant, a processing stabilizer, and a heat stabilizer in the proportions indicated in Table 2 were problended (or premixed). The obtained mixture was supplied through a main feed port of a blastil extruder (30 mm diameter) having one vent port, and another mixture containing 5 parts by weight of a

polyacetal resin copolymer particulate and 0.1 part by weight of a carboxylic acid hydrazide was fed through a side feed port of the downstream of the bent port, and these mixturnes were meth-threaded to prepare a palletized composition. From thus obtained pellets, prescribed test pieces were flampicated with an injection modifing machine, and concerning each test piece, the amount of formaldshyde emission from the test piece, and the bleeding property were measured. The results are shown in Table 2 mixtures are considered to the property of the property of the property were measured.

[0136] Comparative Examples 1 to 5

For comparison, a sample prepared without addition of the carboxylic acid hydrazide compound, and samples with addition of either adjible acid dilydrazide for reuric acid hydrazide as abort-chain fatty acid hydrazide were evaluated in the same meaner described above. The results are shown in Table 3.

[0137] The polyacetal resin copolymers, the carboxylic acid hydrazide compounds, the antioxidants, the processing stabilizers, the heat stabilizers, the coloring agents, and the weather (light)-resistant stabilizers used in the Examples and Comparative Examples are as follows.

[0138] 1. Polyacetal copolymer "a"

- (a-1): Polyacetal resin copolymer (melt index = 9 g/10 min.)
- (a-2): Polyacetal resin copolymer (melt index = 27 g/10 min.)

Incidentally, the melt index was a value (g/10 min.) determined under conditions of 190°C and 2169 g, based on ASTM-D1238.

20 [0139] 2. Carnoxylic acid hydrazide "b"

- (b-1): 8.12-Eicosagienedioic acid dihydrazide
- (b-2): Eicosanediolc acid dihydrazide
- (b-3): 12-Hydroxystearic acid hydrazide (b-4): Montanic acid hydrazide
 - (b-5): 1,4-Cyclohexanedicarboxylic acid dihydrazide
 - (b-6): Linoleic dimer acid dihydrazide
 - (b-7): Adipic ac dihydrazide
 - (b-8): Lauric as-a hydrazide

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[0140] 3. Antioxidant "c"

- (c-1): Triethylene glycol bis[3-(3-t-butyl-5-methyl-4-hydroxyphenyl)propionate]
- (c-2): Pentaerythritol tetrakis[3-(3,5-di-t-butyl-4-hydroxyphenyl)propionate]
- (c-3): 2,2'-methylenebis(4-methyl-6-t-butylphenol)

[0141] 4. Processing stabilizer "d"

- (d-1): Ethylenebisstearylamide
- (d-2): Montanate [manufactured by Toyo-Petrolite Co., Ltd., "LUZA WAX-EP"]
- (d-3): Polyethylene glycol [molecular weight: 35000]

[0142] 5. Heat stabilizer (metal salt of an organic carboxylic acid, alkaline earth metal sait, basic nitrogen-containing compound) "e"

- (e-1): Calcium 12-hydroxysrearate
- (e-2): Magnesium stearate (e-3): Calcium citrate
- (e-3). Calcium cirate
- (e-4): Magnesium oxide (e-5): Allantoin
- (e-5): Aliantol
- (e-6): Biurea
- (e-7): Nylon 6-66-610

[0143] 6. Coloring agent "f"

(f-1): Carbon black (acetylene black)

[0144] 7. Weather (light)-resistant stabilizer "g"

(g-1): 2-{2'-Hydroxy-3',5'-bis(α,α -dimethylbenzyl) phenyl]benzotriazole (g-2): Bis(2,2,6,6-tetramethyl-4-piperidyl)sebacate

[0145] [Table 1]

Table 1

DOGID: 46P_____1674628A1_1_>

							Examples	s					
	1	2	۳	4	2	9	-	80	6	10	=	12	13
Polyacetal resin copolymer	a-1	a-1	a-2	a-1	a-1	8-1	1-8	1-e	- N	1-6	1	1	-
(parts by weight)	100	100	100	100	100	100	100	100	100	100	100	100	100
Carboxylic acid hydrazide	1	-	1	1		-	1	1					
• 0	1	1	1-0	1-0	T 0	7-0	1-0	7-0	D-3	0-3	9-0	D-5	9-q
(parts by weight)	0.5	0.3	0.3	0.2	0.2	0.3	0.3	0.2	0.3	0.2	0.2	0.2	0.2
Antioxidant "c"		c-1	c-1	6-1	2-2	6-3	c-3	6-1	6-3	6-3	0-1	0-3	0-1
s by weight)		0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0,3	0.3	0.3
PL ng stabilizer "d"		q-1	q-1	d-2	4-1	d-1	q-3	q-1	d-2	d-2	q-1	d-1	g-1
(parts by weight)		0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.5	0.2	0.5	0.2	0.2
Heat stabilizer "e"		e-1	e-1	e-1	e-2	6-3	e	6-3	0-1	6-3	6-1	6-3	9-1
(parts by weight)		0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03
Coloring agent "f"													
(parts by weight)			,					,		ı	,		,
Weather (light)-resistant								-	-		-		
stabilizer "g"	,	,			,	,	,	1	,	,	,		
(parts by weight)													
Moldability (Mold deposit)	~	4	4	2	2	2	S	2	5	2	ıc	4	u
Amount of formaldehyde			1		1	1		1	1				
emission, Dry (µg/cm2)	0.04	0.03	6.03	0.03	0.05	0.03	0.04	90.0	0.04	0.04	0.02	0.04	0.05
Amount of formaldehyde	000	000				1	1	1	:	1			
emission, Humid (µg/cm2)	0.00	0.08	90.0	0.07	0.08	0.02	0.07	0.08	0.00	0.06	0.08	0.01	0.08
Bleeding property	В	A	A	A	P	4	q	d	4	d	-	4	-

195000E «EP_____1674526A1_/_>

Table 2

			Examples	103		
	14	15	16	17	18	19
Polyacetal resin copolymer	a-1	a-1	a-1	a-1	a-1	1
(parts by weight)	100	100	100	100	100	100
Carboxylic acid hydrazide	1	F-1	-	- 4		
a.		1	7-0	7-0	1-0	r-0
(parts by weight)	0.1	0.1	0.1	0.3	0.3	0.3
Antioxidant "c"	c-1	c-1	0-1	0-1	0-1	6
(parts by weight)	0.3	0.3	0.3	0.3	0.03	0 3
Processing stabilizer "d"	d-1	g-1	d-2	q-1	d-1	-P
(parts by weight)	0.5	0.3	0.2	0.2	0.2	0.2
Heat stabilizer "e"	e-4 e-5	e-4 e-6	e-3 e-7	e-1	e-1	9
(parts by weight)	0.03 0.05	0.03 0.05	0.03 0.05	0.1	0.1	
Coloring agent "f"				£-1		4
(parts by weight)				0.5		1 6
Weather (light)-resistant						
stabilizer "g"		ı			g-1 g-2	٠
(parts by weight)					0.4 0.2	
Moldability (Mold deposit)	4	4	4	4	4	4
Amount of formaldehyde						
emission, Dry (µg/cm2)	0.03	0.03	0.04	0.13	0.22	0.11
Amount of formaldehyde						
emission, Humid (µg/cm2)	0.02	0.04	90.0	0.15	0.23	0.13
Bleeding property	A	A	A	B	B	m

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		Compar	Comparative Examples	xamples	,,,
	1	2		4	2
olyacetal resin copolymer "a"	a-1	a-1	a-1	a-1	a-1
(parts by weight)	100	100	100	100	1.00
Carboxylic acid hydrazide "b"		p-7	p-8	p-7	9-q
(parts by weight)		0.5	0.5	0.3	0.3
Antioxidant "c"	0-1			C-1	6-1
(parts by weight)	0.3		1	0.3	0.3
Processing stabilizer "d"	q-1			d-1	g-1
(parts by weight)	0.2			0.2	0.2
Heat stabilizer "e"	e-1			e-1	6-1
(parts by weight)	0.1			0.03	0.03
Coloring agent "f"					
(parts by weight)	ı				
Weather (light)-resistant					
stabilizer "g"	1	т	1	1	٠
(parts by weight)					
Moldability (Mold deposit)	S	-	п	-	1
Amount of formaldehyde	00,	3		100	
emission, Dry (µg/cm²)	4.00	0.0	0.00	0.00	0.0
Amount of formaldehyde	03.1	0	30 0	9	9
emission, Humid (µg/cm2)	1.30	0.0	0.00	0.10	
Bleeding property	×	υ	U	U	O

It is apparent from the Tables that as compared with the amount of formaldehyde emission from the resin compositions of Comparative Examples, that from the resin compositions according to Examples is considerably decreased or reduced, and the working and using environment can be drastically improved. Further, in Examples, moldability (mold deposit) is improved, and bleading out is inhibited, thereby the quality of the shaped artials is improved.

Claims

- 1. A polyacetal resin composition comprising a polyacetal resin and a carboxylic acid hydrazide, wherein the carboxylic acid hydrazide comprises at least one member selected " " " group consisting of a saturated or unsaturated or carboxylic acid hydrazide, a saturated or " ununated alicyclic carboxylic acid hydrazide, a saturated or timer acid chydrazide, and an oxycarboxylic acid hydrazide, a dimer acid chydrazide, and an oxycarboxylic acid hydrazide corresponding to each of said hydrazides.
- 2. A resin composition according to claim 1, wherein the carboxylic acid hydrazide comprises at least one member selected from the group consisting of a saturated or unsaturated alliphatic C₁₆₋₄₀ achoxylic acid hydrazide, a saturated or unsaturated insent C₂₀₋₄₀ diffurer acid hydrazide, a saturated or unsaturated insent C₂₀₋₄₀ diffurer acid hydrazide, a saturated or unsaturated insent C₂₀₋₄₀ diffurer acid hydrazide, a saturated or unsaturated insent c₂₀₋₄₀ diffurer acid hydrazide, a saturated or unsaturated cincip cacify hydrazide constitutions.

responding to each of said hydrazides.

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- 3. A resin composition according to claim 1, wherein the carboxytic acid hydrazdic comprises at least one member selected from the group consisting of a seturated crimated alliphatic C₁₆₋₄₀monocarboxytic acid monohydrazdic, a saturated or unsaturated alliphatic C₁₆₋₄₀dicarboxytic acid monohydrazdic, a saturated alliphatic oxy-C₁₆₋₄₀dicarboxytic acid monohydrazdic, a saturated alliphatic oxy-C₁₆₋₄₀dicarboxytic acid monohydrazdic, a saturated or unsaturated alliphatic oxy-C₁₆₋₄₀dicarboxytic acid monohydrazdic, a saturated or unsaturated alliphatic oxy-C₁₆₋₄₀dicarboxytic acid monohydrazdic, a saturated or unsaturated alliphatic or unsaturated alliphatic C₂₆₋₂₀dicarboxytic acid monoh or dihydrazdic, a saturated or unsaturated or unsa
 - 4. A resin composition according to cleim 1, wherein the carboxylic acid hydrazide comprises at least one member selected from the group consisting of montanic acid hydrazide, eicosanediolc acid dihydrazide, 8,12-eicosacilenediolc acid dihydrazide, 12-hydroxystearic acid hydrazide, 1,4-cyclohexanedicarboxylic acid dihydrazide, and linoleic dimer acid dihydrazide.
 - A resin composition according to claim 1, wherein the proportion of the carboxylic acid hydrazide is 0.001 to 20 parts by weight relative to 100 parts by weight of the polyacetal resin.
 - A resin composition according to claim 1, which further comprises at least one member selected from the group consisting of an antioxidiant, a heat stabilizer, a processing stabilizer, a weather (fight)-resistant stabilizer, an impact resistance improver, a slip-improving agent, a coloning agent, and a filter.
- A resin composition according to claim 6, wherein the antioxident, the processing stabilizer, the heat stabilizer, and
 the weather (light)-resistant stabilizer are substantially free from an intramolecular ester bond.
 - A resin composition according to claim 6, wherein the antioxidant comprises at least one member selected from the group consisting of a hindered phenol-series compound and a hindered amine-series compound,
 - A resin composition according to claim 6, wherein the processing stabilizer comprises at least one member selected from the group consisting of a long-chain fatty acid or a derivative thereof, a polyoxyalkylene glycol, and a siliconeseries compound.
- 35 10. A resin composition according to claim 6, wherein the heat stabilizer comprises at least one member selected from the group consisting of a basic introgen-containing compound, a phosphine-series compound, a metal sat of an organic carboxylic acid, an alkalif or islainine earth metal compound, a hydrotalicia, and a zeolite.
 - 11. A resin composition according to claim 6, wherein the heat stabilizer comprises at least one member selected from the group consisting of an alkaline earth metal salt of an organic carboxylic acid, and an alkaline earth metal oxide.
 - A resin composition according to claim 6, wherein the heat stabilizer comprises an alkaline earth metal salt of an oxy-acid.
- 45 13. A resin composition according to claim 6, wherein the weather (light)-resistant stabilizer comprises at least one member selected from the group consisting of a benzorirazole-series compound, a benzophenone-series compound, an aromatic benzoate-series compound, a cyanoacrylate-series compound, a oxalic anillide-series compound, and a hydroxyaryl-1,3,5-flazine-series compound.
- 50 14. A resin composition according to claim 6, wherein the impact resistance improver comprises at least one member selected from the group consisting of a thermoplastic polyurethane and an acrylic core-shell polymer.
 - 15. A resin composition according to claim 6, wherein the slip-improving agent comprises at least one member selected from the group consisting of an olefinic polymer, a silicone-series resin, and afluorine-containing resin.
 - 16. A process for producing a polyacetal resin composition, which comprises mixing a polyacetal resin with a carboxylic acid hydrazide comprising at least one member selected from the group consisting of a saturated or unsaturated long-chial reliphatic carboxylic acid hydrazide, and, a saturated or unsaturated alloyic carboxylic acid hydrazide, and

an oxycarboxylic acid hydrazide corresponding to each of said hydrazides, wherein the resin composition is prepared by using an extruder, and feeding at least said carboxylic acid hydrazide through a side feed port of the extruder,

- 17. A shaped article formed from a polyacetal resin composition recited in claim 1.
- 18. A shaped article according to claim 17, wherein (1) the emission of formatidehyde from the shaped article which is maintained in a closed space for 24 hours at a temperature of 80°C is not more than 1.0 µg per one cm² of the surface area of the article, and/or (2) the emission of formatidehyde from the shaped article which is maintained in a closed space for 3 hours at a temperature of 80°C under saturated humidity is not more than 2 µg per one cm² of the surface area of the article.
- 19. A shaped article according to claim 17, which is an automotive part, an electric or electronic device part, an architectural or pipelline part, a household utensil or cosmetic article part, or a medical device part.

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	INTERNATIONAL SEARCH REPORT	1	International applic	ation No.
			PCT/JP2	004/013825
A. CLASSIP Int.C.	CATION OF SUBJECT MATTER 7 C08L59/00, C08K5/25, C08L71/	02, CD8L83/04		
According to It	aternational Patent Classification (IPC) or to both nation	al classification and IPC	:	
B. FIELDS S	EARCHED			
	mentation searched (classification system followed by c ? COBI, COBK	lassification symbols)		
	searched other than minimum documentation to the ext			
ENECTROPIC DATA	hase consulted during the international search (name of	data base and, where pr	acticable, scarch ten	ns used)
C. DOCUME	NTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where ap			Relevant to claim No.
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X Further do	cuments are listed in the continuation of Box C.	See patent fami	ly annex.	
A* dooument of to be of pure of the pure o	upries of our decomment: feling the giorent size of the an which is not comindent closular relativation. In the property of	"I" laser document pubdet and as in som the principle or thetale on the continue of parties considered asvet of step when the principle of the document of parties considered as vet of the continue of parties of the continue with one being obtained to the continue of the continu	lished after the incomplication of the thing of the most under relevance; the class consistence is the classification of the consistence in the classification of the classifica	med invention cannot be d to involve an inventive med invention cannot be when the decement is summits, such combination by
Japanes	se Patent Office			
	0 (second sheet) (January 2004)	Telephone No.		

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International application No. PCT/JP2004/013825

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